| | | | NT | AA | | |
|--|--|---|---|--|---|---|
| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
| A17503001001_23617338_f1_66 | 3435 | 7207 | 528 | 175 | 215 | 1.2e-24 |
| Description | | | | | | |
| sp:[LN:YAGU_ECOLI] [AC:P77262] [DE:HYPOTHETICAL 23.0 KD PROTE [DB:swissprot] >pir:[LN:G64754 yagU] [GN:yagU] [OR:Escherich [LN:ECU73857] [AC:U73857] [OR: [DE:Escherichia coli chromosom [LE:10789] [RE:11403] [DI:dire [AC:AE000136:U00096] [PN:orf, I Unknown] [OR:Escherichia coli] MG1655 section 26 of 400 of the to 46 residues of approx.] [LE | IN I | TTF-EAEH [4754] [DB:pi chia col es 6-8.] [GI:g17 cical pre apept-bc etegenome | INTERC [PN:pro r2] >g] i] [DB [NT:h] 86481] otein] t2] [DI e.] [N | GENIC obable o:[GI: :genpe ypothe [LN:A [GN:y E:Esch | REGION] membran g1657488 pt-bct1] tical pr E000136] agU] [FN erichia ; 26 pct | ne protein cotein] forf; coli K-12 |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_23621010_c1_971 | 3436 | 7208 | 843 | 280 | 167 | 1.5e-10 |
| Description | 1 | | | | | |
| pir: [LN:E69777] [AC:E69777] [Nomble of the pir: [LN:E69777] [AC:E69777] [Nomble of the pir: [N:E69777] [AC:E69777] [Nomble of the pir: [N:E69777] [N:E69777] [Nomble of the pir: [N:E69777] [| acillus :AB00148 illus su subtili gree.] [E:95608] :BSUB000 ilis] [D | subtilia [AC:2] Subtilia | s] [DB AB00148 (strain e seque ABLE H mplemen Z99106 pt-bct 1 to61 | :pir2] 38] [G 1:168) ence, TH_ARA 1t] :AL009 1] [DE 1850.] | N:ydeC] DNA] 148 kb s C_FAMILY 126] [GN :Bacillu [NT:sim | sequence of OF [:ydeC] ss subtilis |
| ORF Name AI7503001001_23626000_f2_582 Description | NT ID | <u>AA ID</u> 7209 | NT LN 168 | <u>AA</u> <u>LN</u> 55 | <u>Score</u> | <u>P-Value</u> |
| NO-HIT | | | | | | |
| ORF Name AI7503001001_23629040_c2_1318 Description | NT ID | AA ID | NT LN 132 | AA LN 43 | Score | P-Value |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|---|---|--|--|---|---|-------------------------------|
| AI7503001001 23631928 £2 520 | 3439 | 7211 | 228 | <u>الله</u> 75] [| 7 | |
| Description | |] | | J L | | |
| NO-HIT | | | | | | |
| | | _ | | | | |
| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | P-Value |
| A17503001001_23634651_c1_964 | 3440 | 7212 | 1044 | 347 | 549 | 5.0e-53 |
| Description | | | | | | |
| <pre>>gp:[GI:e1185912:g2635523] [LN [FN:unknown] [OR:Bacillus subt complete genome (section 16 of two-component sensor histidine [DI:complement] >gp:[GI:g22931 transduction protein kinase] [DB:genpept-bct2] [DE:Bacillus [LE:65922] [RE:66926] [DI:direction</pre> | ilis] [E 21): fr kinase] 76] [LN: GN:ytsB] subtili | B:genpe om 2997 [LE:11] AF008220 | pt-bct 771to 3501] 0] [AC cillus | 1] [DE 321341 [RE:11 :AF008 subti | :Bacillu 0.] [NT: 4505] 220] [PN lis] | subtilis similar to |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_23672137_c2_1196 | 3441 | 7213 | 957 | 318 | 260 | 6.6e-32 |
| Description | | | | | | |
| <pre>pir:[LN:S76964] [AC:S76964] [sp.] [SR:PCC 6803, , PCC 6803] >gp:[GI:d1019609:g1653966] [LN protein] [OR:Synechocystis sp. [DB:genpept-bct1] [DE:Synechoc 3418852-3573470.] [NT:ORF_ID:S</pre> | [SR:PCC:D90917] [SR:Syystis sp | 6803,] [AC:D90 nechocys . PCC680 | DB: _] [DB: _] [DB: _] [DB: _] [DB:] [DB:] [DB:] | pir2] B00133 p. (st plete | 9] [PN:4 rain:PCC genome, | 7 kD 26803) DNA] 27/27, |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503001001_23672562_c1_924 | 3442 | 7214 | 297 | 98 | 86 | 0.0034 |
| Description | | | | | | |
| <pre>gp:[GI:e8900:g1335718] [LN:PFR] eryrthrocyte surface antigen] [SR:malaria parasite P. falcipa Ag46 RESA mRNA for ring-infector [LE:<1] [RE:>955] [DI:direct]</pre> | [GN:RESA arum] [D |] [OR:P] B:genper | lasmod: pt-inv: | ium fa 1] [DE | lciparum :P.falci | 1] |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|--|---|--------------------------------------|--------------------------------------|--|-----------------------------------|
| A17503001001_23703750_c1_1095 | 3443 | 7215 | 639 | 212 | 7 | |
| Description | | | | l L | - - | |
| NO-HIT | | | | | | |
| *** | | - | | | · | |
| ORF Name | NT ID | AA ID | <u>NT</u> <u>LN</u> | AA LN | Score | P-Value |
| AI7503001001_23728412_c3_1509 | 3444 | 7216 | 699 | 232 | 726 | 8.7e-72 |
| Description | | | | | | |
| pir:[LN:C70020] [AC:C70020] [[GN:yusB] [CL:probable transp [DB:pir2] >gp:[GI:e1184352:g26 [GN:yusB] [FN:unknown] [OR:Bac subtilis complete genome (sect [NT:similar to hypothetical pr [DI:complement] | ort prot 35770] (illus su ion 17 c | cein yae [LN:BSUB abtilis] of 21): | E] [OR 0017] [DB:ge from 3: | :Bacil [AC:Z9 enpept 197001 | lus subt 9120:ALC -bct1] (to 34144 | cilis] 009126] [DE:Bacillus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_23844575_c2_1274 | 3445 | 7217 | 1062 | 353 | 1442 | 1.2e-147 |
| Description | | | | | | |
| <pre>gp:[GI:e1393153:g4490612] [LN: [GN:gapR] [OR:Staphylococcus a aureus gap operon (gapR, gap, [DI:direct]</pre> | ureus] [| DB:genp | ept-bc | t1] [D | E:Staphy | lococcus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_23984775_f1_83 | 3446 | 7218 | 153 | 50 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | P-Value |
| AI7503001001_23989061_f2_399 | 3447 | 7219 | 138 | 45 | ר | |
| Description | | · — — | | | _ | |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|---------------------------------|---------------------------------|-------------------------------|---------------------------|---------------------------------|----------|
| AI7503001001_2401430_c3_1473 | 3448 | 7220 | 762 | 253 | 1076 | 7.1e-109 |
| Description | | | | | | |
| <pre>gp:[GI:e1387399:g4379428] [LN: reductase] [GN:trxB] [OR:Staph; [EC:1.6.4.5] [DE:Staphylococcu [DI:direct]</pre> | ylococcu | s aureus | s] [DB | genpe: | pt-bct1] | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24022582_f2_316 | 3449 | 7221 | 150 | 49 | | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24033567_c2_1316 | 3450 | 7222 | 450 | 149 | 207 | 8.6e-17 |
| Description | | | | | | |
| pir:[LN:G70023] [AC:G70023] [[OR:Bacillus subtilis] [DB:pir [AC:Z99120:AL009126] [GN:yutE] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [LE:1 | 2] >gp:[[FN:unk subtili | GI:e1184 nown] [G | 1309:g2 DR:Baci ete ger | 263572 illus nome (| 7] [LN:E subtilis section | SUB0017] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24218785_c2_1275 | 3451 | 7223 | 1047 | 348 | 1597 | 4.4e-164 |
| Description | | | | | | |
| gp:[GI:e1393154:g4490613] [LN: [PN:glyceraldehyde-3-phosphate binding protein] [OR:Staphyloc [DE:Staphylococcus aureus gap [NT:putative] [LE:1845] [RE:28 | dehydro occus au operon (| genase] reus] [I gapR, ga | [GN:ga DB:geng | ap] [F pept-b | ct1] | , |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24230001_f2_362 | 3452 | 7224 | 258 | 85 | | |
| <u>Description</u> | | | | | | |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|---------------------|---------------------|-------------------|------------------|-------------------|----------------|
| AI7503001001_24240888_f1_281 | 3453 | 7225 | 129 | 42 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| | | | NT | 7.7 | · - · | |
| ORF Name | NT ID | AA ID | LN | <u>AA</u> LN | Score | <u>P-Value</u> |
| A17503001001_24242285_c3_1516 | 3454 | 7226 | 1056 | 351 | 439 | 2.2e-41 |
| Description | | - | | | | |
| <pre>gp:[GI:g4981378] [LN:AE001751] protein] [GN:TM0845] [OR:Thermo [DE:Thermotoga maritima section [NT:similar to GB:AE000783 pero [DI:direct]</pre> | otoga ma n 63 of | ritima] 136 of | [DB:ge the com | npept plete | -bct2] genome. |] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AT7503001001_24251500_f2_416 | 3455 | 7227 | 135 | 44 | 7 | |
| Description | | | | . | - | |
| NO-HIT | | | - | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503001001_24253153_f1_150 | 3456 | 7228 | 129 | 42 |] | |
| Description | - | • | • | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_24256287_c2_1260 | 3457 | 7229 | 141 | 46 |] | |
| <u>Description</u> | | | | | | |
| NO-HIT | | | | | | <u> </u> |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_24258388_c3_1405 | 3458 | 7230 | 507 | 168 | 102 | 0.0053 |
| <u>Description</u> | | | | | | |
| <pre>gp:[GI:e1407794:g4493938] [LN:E [OR:Plasmodium falciparum] [SR: [DB:genpept-inv1] [DE:Plasmodium] [NT:predicted using hexexon; MA]</pre> | malaria ım falci | . parasi parum M | te P. f AL3P5, | alcipa comple | ete sequ | ence.] |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|--|--|--|---|---|---------------------------------|------------------------------------|
| A17503001001_24259762_f3_679 | 3459 | 7231 | 132 | 43 | ٦ | |
| Description | —————————————————————————————————————— | | · | | _ | |
| NO-HIT | | | | | | |
| | | | | | | |
| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | <u>AA</u> LN | Score | <u>P-Value</u> |
| A17503001001_24266888_c3_1413 | 3460 | 7232 | 123 | 40 | 7 | |
| Description | | <u> </u> | | | _ | |
| NO-HIT | | | | | | |
| | | | NT | <u>AA</u> | | |
| ORF Name | NT ID | AA ID | LN | LN | Score | <u>P-Value</u> |
| AI7503001001_24272125_c2_1292 | 3461 | 7233 | 261 | 86 | 72 | 0.024 |
| Description | | | | | | |
| <pre>gp:[GI:g942589] [LN:MIU29676] iowae] [DB:genpept-bct2] [DE:M sequence, and 23S rRNAgene, pa</pre> | lycoplasm | na iowae | 16S rF | NA ge | ne, comp | plete |
| | | | | | | |
| ORF Name | NT ID | AA ID | $\frac{\text{NT}}{\text{LN}}$ | <u>AA</u> LN | Score | P-Value |
| ORF Name AI7503001001_24297000_f2_548 | NT ID 3462 | | | | Score | <u>P-Value</u> |
| | | | LN | LN | Score | <u>P-Value</u> |
| AI7503001001_24297000_f2_548 | | | LN | LN | Score | P-Value |
| AI7503001001_24297000_f2_548 Description | | | LN | LN | Score | P-Value |
| Description NO-HIT | 3462 | 7234 AA ID | <u>LN</u> 225 . | <u>LN</u> 74 |] | |
| Description NO-HIT ORF Name | 3462 NT ID | 7234 AA ID | LN 225 . NT LN | <u>LN</u> 74 <u>AA</u> <u>LN</u> | Score | P-Value |
| Description NO-HIT ORF Name A17503001001_24329053_c1_979 | NT ID 3463] [AC:AH B:genper by C. e] | AA ID 7235 7078790] ot-inv2] egans cl | LN 225 NT LN 444 [GN:F3 [DE:Ca | AA LN 147 147 166H12. Renorh | Score 142 3] abditis | P-Value 2.8e-09 elegans |
| Description NO-HIT ORF Name A17503001001_24329053_c1_979 Description gp:[GI:g3329623] [LN:CELF36H12 [OR:Caenorhabditis elegans] [D cosmid F36H12.] [NT:coded for | NT ID 3463] [AC:AH B:genper by C. e] | AA ID 7235 7078790] ot-inv2] egans cl | LN 225 NT LN 444 [GN:F3 [DE:Ca | AA LN 147 147 166H12. Renorh | Score 142 3] abditis | P-Value 2.8e-09 elegans |
| Description NO-HIT ORF Name Al7503001001_24329053_c1_979 Description gp:[GI:g3329623] [LN:CELF36H12 [OR:Caenorhabditis elegans] [D cosmid F36H12.] [NT:coded for [LE:21606:21872:22500:22677] [| NT ID 3463] [AC:AH B:genper by C. el RE:21701 | AA ID 7235 7078790] pt-inv2] egans cE 1:22454:2 | LN 2225 . NT LN 444 [GN:F3 [DE:Ca DNA CEM 22629:2 | AA LN 147 36H12. aenorh 1SF30F 22875] | Score 142 3] abditis [DI:dir | P-Value 2.8e-09 elegans rectJoin |
| Description NO-HIT ORF Name AT7503001001_24329053_c1_979 Description gp:[GI:g3329623] [LN:CELF36H12 [OR:Caenorhabditis elegans] [D cosmid F36H12.] [NT:coded for [LE:21606:21872:22500:22677] [ORF Name | NT ID 3463] [AC:AF B:genper by C. el RE:21701 | AA ID 7235 7078790] pt-inv2] egans cE 1:22454:2 | LN 225 . NT LN 444 [GN:F3 [DE:Ca DNA CEM 22629:2 | AA LN 147 Aenorh ISF30F 2875] | Score 142 3] abditis [DI:dir | P-Value 2.8e-09 elegans rectJoin |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
|---|--|---|---|---|--|--|
| AI7503001001_24345257_f3_652 | 3465 | 7237 | 168 | 55 | 7 | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24347175_c3_1416 | 3466 | 7238 | 165 | 54 |] | |
| Description | | | | | | |
| NO-HIT | | | | | _ | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_24391937_c1_1068 | 3467 | 7239 | 759 | 252 | 105 | 0.0062 |
| [CL:tetratricopeptide repeat h >gp:[GI:e1186169:g2635994] [LN [FN:unknown] [OR:Bacillus subt complete genome (section 18 of [RE:177129] [DI:complement] >g [AC:Z94043] [PN:hypothetical p [DB:genpept-bct1] [DE:B.subtil [RE:5996] [DI:direct] | :BSUB001 ilis] [I 21): fr p:[GI:e3 rotein] | .8] [AC: DB:genper rom 3399 13021:g [GN:yvcl | Z99121 pt-bct: 551to : 194564! D] [OR | :AL0091 1] [DE 3609060 5] [LN :Bacil1 | 126] [GN :Bacillu D.] [LE: :BSZ9404 lus subt | I:yvcD] us subtilis 175675] us:ilis] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24397952_c1_1067 | 3468 | 7240 | 681 | 226 | 119 | 0.00011 |
| Description | | | | | | |
| <pre>pir:[LN:E70031] [AC:E70031] [[CL:tetratricopeptide repeat h >gp:[GI:e1186169:g2635994] [LN [FN:unknown] [OR:Bacillus subt complete genome (section 18 of [RE:177129] [DI:complement] >g [AC:Z94043] [PN:hypothetical p [DB:genpept-bct1] [DE:B.subtil [RE:5996] [DI:direct]</pre> | omology] :BSUB001 ilis] [E 21): fr p:[GI:e3 rotein] | [OR:Bac .8] [AC: .8:genpe] .50m 33999 .13021:gi | cillus Z99121: pt-bct: 551to 3 194564! D] [OR: | subtil :AL009: L] [DE: 3609060 5] [LN: :Bacil] | lis] [DE 126] [GN :Bacillu D.] [LE: :BSZ9404 lus subt | s:pir2] I:yvcD] s subtilis 175675] 3] ilis] |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
|---|--|---|--|--|--|--|
| A17503001001_24406577_c3_1415 | 3469 | 7241 | 390 | 129 | 7 | |
| Description | | · | | · · · · · · · · · · · · · · · · · · · | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | $\frac{\text{NT}}{\text{LN}}$ | AA LN | Score | P-Value |
| AI7503001001_24407637_f1_57 | 3470 | 7242 | 243 | 80 | | • |
| Description | | | | | | |
| NO-HIT | | · | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_24415930_f2_310 | 3471 | 7243 | 1068 | 355 | 977 | 2.2e-98 |
| [AC:Z99120:AL009126] [GN:yutJ] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:st [RE:111916] [DI:direct] | subtil | is comple | te ge | nome (| section | 17 of 21): |
| ORF Name A17503001001 24429650 c3 1408 | NT ID | AA ID | <u>NT</u> <u>LN</u> 1197 | <u>AA</u> <u>LN</u> [398 | <u>Score</u> | P-Value 1.0e-84 |
| Description | 3772 |]/244 | 1177 | 398 | المين | 1.06-04 |
| sp:[LN:NAGA_BACSU] [AC:O34450] [EC:3.5.1.25] [DE:DEACETYLASE)] [AC:A69664] [PN:N-acetylglucos] [OR:Bacillus subtilis] [DB:ps [AC:Z99121:AL009126] [PN:N-acetylglucosar [GN:nagA] [FN:N-acetyl glucosar [DB:genpept-bct1] [EC:3.5.1.25] (section 18 of 21): from 339955 [RE:196056] [DI:direct] >gp:[GS [PN:N-acetylglucosamine 6-P des [DB:genpept-bct2] [DE:Bacillus [LE:31777] [RE:32967] [DI:compo | [SP:03 samine-6 ir2] >gr tylgluco mine uti [DE:Ba 51to 360 I:g26188 acetylas subtili | 34450] [D 5-phospha p:[GI:e11 psamine-6 ilization acillus s 29060.] [356] [LN: se] [GN:n | B:swiste dea 86189 -phosp] [OR ubtil: SP:034 AF0173 | ssprot acetyla g2636 phate of Bacila is comp 4450] L13] [2 [OR:Bac |] >pir:[ase nagA 014] [LN deacetyl lus subt plete ge [LE:1948 AC:AF017 cillus s | [GN:nagA I:BSUB0018] [ase] [ilis] [nome [66] [113] [subtilis] |

NT AAORF Name NT ID AA ID Score P-Value LN LN AI7503001001_24479842 c1 1106 3473 7245 558 185 256 5.5e-22 Description pir:[LN:D69800] [AC:D69800] [PN:conserved hypothetical protein yfhC] [GN:yfhC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182838:g2633172] [LN:BSUB0005] [AC:Z99108:AL009126] [GN:yfhC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 5 of 21): from 802821 to1011250.] [NT:similar to hypothetical proteins] [LE:120274] [RE:120858] [DI:direct] >gp:[GI:d1025385:g2804533] [LN:D85082] [AC:D85082] [PN:YfhC] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, genome sequence, 79 to 81 degree region.] [LE:13165] [RE:13749] [DI:direct] NT AAORF Name NT ID AA ID Score P-Value LN LN AI7503001001_24490676_c2_1179 141 3474 7246 46 Description NO-HIT NTAΑ ORF Name NT ID AA ID Score P-Value LN LNAI7503001001_24506712_c2_1240 3475 7247 324 5.3e-88 Description gp:[GI:e1330453:g3724156] [LN:SAA005352] [AC:AJ005352] [PN:membrane protein] [GN:sstB] [FN:iron transport] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus, Sst putative iron transport operon.] [LE:1451] [RE:2132] [DI:direct] NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001_24507932_c1_1027 3476 7248 199 5.6e-29 600 322 Description pir:[LN:D69924] [AC:D69924] [PN:hypothetical protein yorS] [GN:yorS] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185499:g2634420] [LN:BSUB0011] [AC:Z99114:AL009126] [GN:yorS] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [LE:172005] [RE:172523] [DI:complement] >gp:[GI:g3025618] [LN:AF020713] [AC:AF020713] [PN:unknown] [GN:yors] [OR:Bacteriophage SPBc2] [DB:genpept-phg] [DE:Bacteriophage SPBc2 complete

genome.] [LE:112997] [RE:113515] [DI:direct]

| ORF Name | NT ID | AA ID | $\frac{\mathtt{NT}}{\mathtt{LN}}$ | AA LN | Score | P-Value |
|---|--|--|---|--|--|--|
| A17503001001_24611512_c2_1139 | 3477 | 7249 | <u>—</u> 129 | 42 | 7 | |
| Description | | ······································ | | | - | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24616050_f2_445 | 3478 | 7250 | 150 | 49 |] | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_24640828_c2_1138 | 3479 | 7251 | 516 | 171 |] | |
| <u>Description</u> | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_24642632_f3_765 | 3480 | 7252 | 1245 | 414 | 1245 | 8.7e-127 |
| Description | | | | | | |
| sp:[LN:PEPT_BACSU] [AC:P55179] [EC:3.4.11] [DE:PEPTIDASE T, [SP:P55179] [DB:swissprot] >pir [PN:aminotripeptidase,:peptidase [EC:3.4.11] [DB:pir2] >gp:[GI [GN:pepT] [FN:unknown] [OR:Baci [DE:B.subtilis orfs 1,2,3,4, per [RE:6499] [DI:direct] >gp:[GI:er [AC:Z99123:AL009126] [PN:peptidase [BE:196699] [DI:direct] >gp:[GI:er [RE:196699] [DI:direct] >gp:[GI [AC:D83026:D45911] [GN:pepT] [GI (strain:BGSC 1A1) DNA] [DB:genpare sequence covering lic-cel region (peptidase T)] [LE:27766] [RE:2 | (AMINOTO INTERPRETATION OF THE PROPERTY OF THE | FRIPEPTII 59674] [F 59674] [F 59674] [F 59:914292 ubtilis] galE ger L:g263642 (tripepti L-] [DE: 01to 4010 379:g1783 ulus subt c1] [DE:F | DASE) AC:H696 [OR:I 259] [1 [DB:genes.] 27] [LI 1dase) [2550.] 3235] 31113 homolo | (TRIPE 574] Bacill LN:BSG Enpept [SP:P5 LUS SUB [SP:P [LN:D8 [SR:Bassub ESR:Bassub | PTIDASE) us subti ALE] [AC -bct1] 5179] [I 0020] pepT] [C btilis c 55179] [3026] acillus tilis ge | lis] 2:X99339] E:5267] OR:Bacillus complete [LE:195467] subtilis cnome |

NT AΑ NT ID ORF Name AA ID Score P-Value LN 109 AI7503001001_24650427_f2_349 3481 7253 330 331 6.2e-30 Description pir: [LN:C69772] [AC:C69772] [PN:thioredoxin homolog ydbP] [GN:ydbP] [CL:thioredoxin:thioredoxin homology] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:d1020045:g1881265] [LN:AB001488] [AC:AB001488] [GN:ydbP] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genome sequence, 148 kb sequence of the regionbetween 35 and 47 degree.] [NT:PROBABLE THIOREDOXIN.] [LE:40816] [RE:41136] [DI:complement] >qp:[GI:e1182421:q2632755] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:ydbP] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:similar to thioredoxin] [LE:104560] [RE:104880] [DI:complement] NT AA ORF Name NT ID AA ID Score P-Value LN LN1515 AI7503001001 24650462 cl 944 3482 7254 504 848 1.0e-84 Description pir:[LN:G70008] [AC:G70008] [PN:NADH dehydrogenase (ubiquinone) homolog yufD] [GN:yufD] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184241:g2635659] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yufD] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420.] [NT:similar to NADH dehydrogenase (ubiquinone)] [LE:51840] [RE:53282] [DI:direct] AΑ ORF Name NT ID AA ID Score P-Value LN LN 1968 655 AI7503001001 24664840 c2 1203 3483 7255 2.3e-174 Description pir:[LN:H69626] [AC:H69626] [PN:PTS fructose-specific enzyme IIBC component fruA] [GN:fruA] [CL:phosphotransferase system enzyme II, fructose-specific:phosphotransferase system mannitol-specific enzyme II factor III homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185030:g2633811] [LN:BSUB0008] [AC:Z99111:AL009126] [PN:phosphotransferase system (PTS)] [GN:fruA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:113871] [RE:115778] [DI:direct] >gp:[GI:g3282125] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:fructose PTS IIABC] [GN:fruA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus

subtilis mobA-nprE gene region.] [NT:similar to fructose-specific PTS system

IIBC] [LE:14359] [RE:16266] [DI:direct]

NO-HIT

| | | | Nitr | 7.7 | | |
|---|--|--|---|--------------------------------------|--|-------------------------------|
| ORF Name | NT ID | AA ID | <u>NT</u> <u>LN</u> | <u>AA</u> LN | Score | P-Value |
| A17503001001_24796927_f1_61 | 3484 | 7256 | 789 | 262 | 325 | 2.7e-29 |
| Description | | | | | | _ |
| <pre>pir:[LN:E69883] [AC:E69883] [I ymaC] [GN:ymaC] [CL:phage-rela subtilis] [DB:pir2] >gp:[GI:e11 [AC:Z99113:AL009126] [GN:ymaC] [DB:genpept-bct1] [DE:Bacillus from 1781201to 2014980.] [NT:si [RE:82197] [DI:direct]</pre> | ited rep .83386:g [FN:unk subtili | licatio 2634111 nown] [s compl | n prote] [LN:] OR:Bac: ete gen | ein] [BSUB00 illus nome (| OR:Bacil 10] subtilis section | llus 3] 10 of 21): |
| ODP W | | | NT | AA | | |
| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
| AI7503001001_24803125_c1_993 | 3485 | 7257 | 978 | 325 | 982 | 6.5e-99 |
| sp:[LN:MDH_BACIS] [AC:Q59202] [DE:MALATE DEHYDROGENASE,] [SP:[AC:S61213] [PN:malate dehydrogenestillus israeli] [EC:1.1.1 [AC:X90527] [PN:malate dehydrogenestillus] [EC:1.1.1.37] [DB:genpept-bct1] [EC:1.1.1.37] gene.] [SP:Q59202] [LE:291] [RE- | Q59202] pgenase,37] [D genase] [DE:B. | [DB:sw] [CL:L B:pir2] [OR:Bac israeli | issprof -lactat >gp:[G illus : DNA fo | t] >pi te deh GI:g96 israel | r:[LN:Se ydrogena 3019] [I i] | 51213] ase] LN:BIDNAMD] |
| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | AA LN | Score | P-Value |
| AI7503001001_24803332_c2_1305 | 3486 | 7258 | 840 | 279 | 712 | 2.6e-70 |
| Description | | | | | | <u> </u> |
| pir:[LN:B70020] [AC:B70020] [F [GN:yusA] [CL:lipoprotein-28] >gp:[GI:e1184351:g2635769] [LN: [FN:unknown] [OR:Bacillus subticomplete genome (section 17 of hypothetical proteins] [LE:1638 | [OR:Bac BSUB001 lis] [D 21): fr | illus s 7] [AC: B:genpe om 3197 | ubtili: Z99120 pt-bct: 001to : | E] [DB:AL009. L] [DE:B41442 | :pir2] 126] [GN :Bacillu 0.] [NT: | N:yusA] us subtilis |
| ORF Name [AI7503001001 2531500 c3 1360 | NT ID | AA ID | NT LN | AA LN | Score | P-Value |

| ORF Name | NT ID | AA ID | TN N.T. | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|-------|------------|------------|-----------------|--------------|----------------|
| AI7503001001_25401377_c1_1045 | 3488 | 7260 | 852 | 283 | 939 | 2.3e-94 |
| Description | | , <u> </u> | | | , | |
| gp:[GI:e1330454:g3724157] [LN: | | | | | | nding |
| <pre>protein] [GN:SstC] [FN:iron tr [DB:genpept-bct1] [DE:Staphylo</pre> | | | | | | ansport |
| operon.] [LE:2186] [RE:2947] [| | | L | | | |
| ORF Name | NT ID | AA ID | NT | AA | Score | P-Value |
| | | | LN | <u>LN</u> | <u>3001e</u> | <u>F-value</u> |
| A17503001001_25401675_c2_1319 | 3489 | 7261 | 174 | 57 |] | |
| Description | | | | | | |
| NO-HIT | | | | | | _ |
| | | | NT | AA | | |
| ORF Name | NT ID | AA ID | LN | LN | Score | <u>P-Value</u> |
| AI7503001001_25415802_c1_1102 | 3490 | 7262 | 126 | 41 | 7 | |
| Description | | , <u></u> | L | | - | |
| NO-HIT | | | | | | |
| | | - · · · |), TO | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_25473782_c3_1498 | 3491 | 7263 | 186 | 61 | 1 | |
| Description | | | | | J | <i>/</i> |
| NO-HIT | | | | | | |
| | | | | * | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_25492312_f1_302 | 3492 | 7264 | 126 | 41 |] | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |

NTAAORF Name NT ID AA ID Score P-Value LN LN AI7503001001 25572180 c1 956 1272 3493 7265 423 895 1.1e-89

Description

sp:[LN:YXJA BACSU] [AC:P42312] [GN:YXJA:N15HR] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 43.7 KD PROTEIN IN KATB 3'REGION] [SP:P42312] [DB:swissprot] >pir:[LN:G70078] [AC:G70078] [PN:pyrimidine nucleoside transport homolog yxjA] [GN:yxjA] [CL:pyrimidine nucleoside transport protein nupC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1186401:g2636437] [LN:BSUB0020] [AC:Z99123:AL009126] [GN:yxjA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:similar to pyrimidine nucleoside transport] [SP:P42312] [LE:206141] [RE:207334] [DI:direct] >gp:[GI:e1184627:g2636448] [LN:BSUB0021] [AC:Z99124:AL009126] [GN:yxjA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 21 of 21): from 3999281to 4214814.] [NT:similar to pyrimidine nucleoside transport] [SP:P42312] [LE:5261] [RE:6454] [DI:direct] >gp: [GI:d1012369:g665999] [LN:D83026] [AC:D83026:D45911] [GN:yxjA] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:BGSC 1A1) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genome sequence covering lic-cel region.] [NT:homologous to pyrimidine nucleoside transport] [LE:17131] [RE:18324] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> <u>LN</u> | Score | P-Value | |
|-------------------------------|-------|-------|----------|------------------------|-------|----------|--|
| AI7503001001_25585891_c2_1218 | 3494 | 7266 | 1980 | 659 | 1949 | 2.2e-201 | |

Description

pir:[LN:D69815] [AC:D69815] [PN:conserved hypothetical protein yfnI]
[GN:yfnI] [CL:Bacillus subtilis probable anion-binding protein yflE]
[OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182705:g2633039] [LN:BSUB0004]
[AC:Z99107:AL009126] [GN:yfnI] [FN:unknown] [OR:Bacillus subtilis]
[DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 4 of 21):
from 600701 to813890.] [NT:alternate gene name: yetP; similar to
hypothetical] [LE:195080] [RE:197041] [DI:direct] >gp:[GI:d1020933:g2116767]
[LN:D86418] [AC:D86418] [PN:YfnI] [OR:Bacillus subtilis] [SR:Bacillus
subtilis (strain:AC327) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genomic
DNA 69-70 degree region, partialsequence.] [LE:16150] [RE:18111]
[DI:complement]

NT AΑ ORF Name AA ID NT ID Score P-Value LN LN AI7503001001 25806300 f1 275 750 249 3495 7267 1270 2.0e-129

Description

gp:[GI:e255626:g1617428] [LN:SEABCTS] [AC:X99127] [PN:ATP binding protein]
[FN:iron repressible ABC transport system] [OR:Staphylococcus epidermidis]
[DB:genpept-bct1] [DE:S.epidermidis gene encoding ABC transport system.]
[LE:41] [RE:787] [DI:direct]

NT AAORF Name NT ID AA ID Score P-Value LN LN A17503001001_26171927_c2_1246 3496 342 7268 113 221 2.8e-18 Description sp:[LN:YTXJ BACSU] [AC:P39914] [GN:YTXJ] [OR:BACILLUS SUBTILIS] [DE:(ORF3)] [SP:P39914] [DB:swissprot] >pir:[LN:S21420] [AC:S21420:F70003:S71003] [PN:general stress protein homolog ytxJ] [GN:ytxJ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g39815] [LN:BSAROAG] [AC:X65945] [GN:orf 2] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis aroA-aroG gene.] [SP:P39914] [LE:105] [RE:431] [DI:direct] >gp:[GI:e1185849:g2635460] [LN:BSUB0016] [AC:Z99119:AL009126] [GN:ytxJ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [NT:alternate gene name: csb40; similar to general] [SP:P39914] [LE:48061] [RE:48387] [DI:complement] >gp:[GI:g2293219] [LN:AF008220] [AC:AF008220] [PN:YtxJ] [GN:ytxJ] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis rrnB-dnaB genomic region.] [LE:132039] [RE:132365] [DI:direct] NTAΑ ORF Name NT ID P-Value AA ID Score LN LN AI7503001001_26176693_f2_482 3497 828 275 7269 685 1.9e-67 Description pir:[LN:A69162] [AC:A69162] [PN:gufA protein homolog MTH473] [GN:MTH473] [CL:gufA protein] [OR:Methanobacterium thermoautotrophicum] [DB:pir2] >gp:[GI:g2621542] [LN:AE000831] [AC:AE000831:AE000666] [PN:conserved protein] [GN:MTH473] [OR:Methanobacterium thermoautotrophicum] [DB:genpept-bct1] [DE:Methanobacterium thermoautotrophicum from bases 404817 to 415582(section 37 of 148) of the complete genome.] [NT:Function Code:14.01 - Unknown, Conserved protein;] [LE:9769] [RE:10548] [DI:direct] ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 2618827 cl 1042 7270 288 0.023 I75

Description

sp:[LN:LUXY VIBFI] [AC:P21578] [GN:LUXY] [OR:VIBRIO FISCHERI] [DE:YELLOW FLUORESCENT PROTEIN (YFP)] [SP:P21578] [DB:swissprot] >pir:[LN:A36037] [AC:A36037:A39946] [PN:yellow fluorescent protein:luxY protein] [GN:luxY] [CL:riboflavin synthase alpha chain] [OR:Vibrio fischeri] [DB:pir2] >gp:[GI:g155235] [LN:VIBLUXY] [AC:M60852] [PN:yellow fluorescent protein] [GN:luxY] [OR:Vibrio fischeri] [SR:V.fischeri (strain Y-1) DNA] [DB:genpept-bct1] [DE:Vibrio fischeri yellow fluorescent protein (luxY) gene, completecds.] [LE:45] [RE:629] [DI:direct]

| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
|--|--|--|--|--|---|---|
| AI7503001001_26190785_c2_1306 | 3499 | 7271 | 1287 | 428 | 1408 | 4.7e-144 |
| Description | | | | | | |
| pir:[LN:F70019] [AC:F70019] [] [CL:nifS protein] [OR:Bacillon >gp:[GI:e1184347:g2635765] [LN [FN:unknown] [OR:Bacillus subtains and complete genome (section 17 of NifS protein homolog] [LE:1595] | us subti :BSUB001 ilis] [I 21): fr | .lis] [D .7] [AC: B:genpe com 3197 | B:pir2 Z99120 pt-bct 001to |] :AL009 1] [DE 341442 | 126] [GN :Bacillu 0.] [NT: | N:yurW] us subtilis |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_26204837_f1_69 | 3500 | 7272 | 159 | 52 | | |
| Description | | | | | | |
| NO-HIT | | | , | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_26211552_c2_1270 | 3501 | 7273 | 984 | 327 | 1034 | 2.0e-104 |
| Description | | | | | | |
| pir:[LN:B70032] [AC:B70032] [IGN:yvcL] [OR:Bacillus subtil: [LN:BSUB0018] [AC:Z99121:AL009: subtilis] [DB:genpept-bct1] [DI:Bs of 21): from 3399551to 36090 [LE:169083] [RE:170033] [DI:contilis] [AC:Z94043] [PN:BSZ94043] [AC:Z94043] [PN:Bsubtilis] [DB:genpept-bct1] [DI:Contilis] [DB:genpept-bct1] | is] [DB: 126] [GN E:Bacill 060.] [N mplement nypothet E:B.subt | pir2] > I:yvcL] us subt IT:simil] >gp:[ical pr ilis ge | gp:[GI [FN:un] ilis co ar to] GI:e31 otein] nomic] | ell86 known] omplet nypoth 3028:g [GN:y | 163:g263 [OR:Bace genome etical p 1945652] vcL] [OR agment (| s5988] cillus c (section proteins] c:Bacillus (88 kb).] |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|---|---|---|--|--|--|---|
| A17503001001_26259638_f1_134 | 3502 | 7274 | 765 | 254 | 176 | 2.2e-11 |
| Description | 1 | | | | | |
| pir: [LN:B70798] [AC:B70798] [CR:Mycobacterium tuberculosis [LN:MTV025] [AC:AL022121:AL123-GN:Rv3737] [OR:Mycobacterium tuberculosis [NT:Rv3737, (MTV025.085), len: [DI:direct] >gp:[GI:e1264597:g1] [PN:hypothetical protein Rv373-GDB:genpept] [DE:Mycobacterium 155/162.] [NT:Rv3737, (MTV025.4-GRE:96385] [DI:direct] |] [DB:pi 456] [PN tubercul H37Rv c 529. Pr 2960161] 7] [GN:R tubercu | r2] >gp I:hypoth .osis] [:omplete :obable [LN:MT :v3737] | :[GI:e1 etical DB:genp genome membrar V025] [OR:Myc | prote prote pept-be; segn ne] [Li [AC:AL cobacte pmplete | 7:g29601 in Rv373 ct1] ment 155 E:94796] 022121:A erium tu e genome | 61] 7] /162.] [RE:96385] L123456] berculosis] ; segment |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_26359805_f1_11 | 3503 | 7275 | 168 | 55 | | • |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_26360663_c3_1530 | 3504 | 7276 | 1308 | 435 | 1039 | 5.9e-105 |
| Description | | | | | | |
| pir: [LN:A70015] [AC:A70015] [CL:NADH dehydrogenase] [OR:Bac >gp: [GI:e1184289:g2635707] [LN [FN:unknown] [OR:Bacillus subtant complete genome (section 17 of NADH dehydrogenase] [LE:102091] >gp: [GI:e311467:g1934829] [LN:] [GN:yumB] [OR:Bacillus subtilis DNA fragment from yumA to yulf [DI:complement] | cillus s :BSUB001 ilis] [C 21): fr] [RE:10 BSZ93939 s] [DB:g | ubtilis 7] [AC: B:genpe om 3197 3311] [C: AC:Z genpept- |] [DB:r Z99120: pt-bct1 001to 3 DI:comr 93939] bct1] [| Dir2] AL009 [DE 41442 [PN:N. [DE:B. | 126] [GN :Bacillu 0.] [NT: t] ADH dehy subtilis | :yumB] s subtilis similar to drogenase] genomic |
| | | | NT | AA | | n 1 |
| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
| A17503001001_26367175_c3_1476 | 3505 | 7277 | 123 | 40 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |

 ORF Name
 NT ID
 AA ID
 NT LN
 AA Score
 P-Value

 AI7503001001_26376077_c1_1113
 3506
 7278
 192
 63
 101
 1.5e-05

Description

pir:[LN:C70063] [AC:C70063] [PN:hypothetical protein ywmG] [GN:ywmG]
[OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e276829:g1648859] [LN:BSATPC]
[AC:Z81356] [PN:unknown] [GN:ywmH] [OR:Bacillus subtilis] [DB:genpept-bct1]
[DE:B.subtilis atpC gene.] [LE:10910] [RE:11098] [DI:direct]
>gp:[GI:e1184573:g2636192] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywmG]
[FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis
complete genome (section 19 of 21): from 3597091to 3809700.] [LE:172069]
[RE:172257] [DI:complement] >gp:[GI:e1184573:g2636192] [LN:BSUB0019]
[AC:Z99122:AL009126] [GN:ywmG] [FN:unknown] [OR:Bacillus subtilis]
[DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [LE:172069] [RE:172257] [DI:complement]

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 A17503001001_26464707_c3_1493
 3507
 7279
 141
 46
 43
 0.039

Description

pir: [LN:S58357] [AC:S66652:S58357] [PN:pepI protein] [GN:pepI] [OR:Staphylococcus epidermidis] [DB:pir2] >gp:[GI:g945016] [LN:SEPEPGNS] [AC:Z49865] [PN:PepI] [GN:pepI] [OR:Staphylococcus epidermidis] [DB:genpept-bct1] [DE:S.epidermidis pepA, pepB, pepC, pepI, pepP and pepT genes.] [LE:2028] [RE:2237] [DI:direct] >gp:[GI:g398082] [LN:STAPEPA] [AC:L23967] [PN:immunity protein] [GN:pepI] [OR:Staphylococcus epidermidis] [SR:Staphylococcus epidermidis DNA] [DB:genpept-bct1] [DE:Staphylococcus epidermidis lantibiotic (pepA) and immunity protein(pepI) gene, complete cds.] [LE:377] [RE:586] [DI:direct]

NTAAORF Name NT ID AA ID Score P-Value LN LN AI7503001001_26578577_c1_1065 1791 3508 7280 596 2276 1.8e-238

Description

sp:[LN:UVRA BACSU] [AC:034863] [GN:UVRA] [OR:BACILLUS SUBTILIS] [DE:EXCINUCLEASE ABC SUBUNIT A] [SP:034863] [DB:swissprot] >pir:[LN:F69729] [AC:F69729] [PN:excinuclease ABC chain A:excision endonuclease ABC,, chain A:uvrA protein] [GN:uvrA] [CL:excinuclease ABC chain A:ATP-binding cassette homology] [OR:Bacillus subtilis] [EC:3.1.-.-] [DB:pir2] >gp:[GI:e1184422:q2636042] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:excinuclease ABC (subunit A)] [GN:uvrA] [FN:excision of ultraviolet light-induced] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:034863] [LE:12034] [RE:14907] [DI:complement] >gp:[GI:g2618842] [LN:AF017113] [AC:AF017113] [PN:excinuclease ABC subunit A] [GN:uvrA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis 300-304 degree genomic sequence.] [LE:15386] [RE:18259] [DI:direct] >gp: [GI:e1184422:q2636042] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:excinuclease ABC (subunit A)] [GN:uvrA] [FN:excision of ultraviolet light-induced] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:O34863] [LE:12034] [RE:14907] [DI:complement]

NTAA ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 26594050 c3 1524 3509 7281 243 80 378 6.5e-35

Description

gp:[GI:d1013748:g1405337] [LN:D86240] [AC:D86240] [PN:D-alanyl carrier protein] [GN:dltC] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (isolate:KAN96) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus gene for unkown function and dlt operon dltA,dltB, dltC and dltD genes,complete cds.] [NT:Sequence homologous to the dltC genes of] [LE:5303] [RE:5539] [DI:direct] >gp:[GI:g4530243] [LN:AF101234] [AC:AF101234] [PN:D-alanine carrier protein DltC] [GN:dltC] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus dltABCD operon, complete sequence; andunknown gene.] [LE:4276] [RE:4512] [DI:direct]

| OPE Namo | NT TO | מא דה | NT | <u>AA</u> | Saoro | D-Walue |
|--|---|---|---|--|--|-------------------------------------|
| ORF Name | NT ID | AA ID | <u>LN</u> | LN | Score | P-Value |
| A17503001001_26735877_c3_1508 | 3510 | 7282 | 402 | 133 | 446 | 4.1e-42 |
| Description | | | | | | |
| pir:[LN:A70021] [AC:A70021] [yusH] [GN:yusH] [CL:glycine clipoyl/biotin-binding homology >gp:[GI:e1184358:g2635776] [LN [FN:unknown] [OR:Bacillus subt complete genome (section 17 of glycine cleavage system protein | leavage] [OR:Ba :BSUB001 ilis] [I 21): fr | system acillus 17] [AC: DB:genpe com 3197 | protein subtil Z99120 pt-bct 001to | n H: is] [Di :AL009: 1] [DE 341442 | B:pir2] 126] [GN :Bacillu 0.] [NT: | I:yusH] s subtilis similar to |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_26753150_£1_147 | 3511 | 7283 | 336 | 111 | 295 | 4.1e-26 |
| Description | | | | • | | |
| sp:[LN:YKI4_YEAST] [AC:P36078] [SR:,BAKER'S YEAST] [DE:HYPOTH REGION] [SP:P36078] [DB:swissp: [PN:hypothetical protein YKL08- [MP:11L] >gp:[GI:g486123] [LN [OR:Saccharomyces cerevisiae] [DE:S.cerevisiae chromosome XI [SP:P36078] [LE:382] [RE:732] | ETICAL 1 rot] >pi 4w] [OR: :SCYKL08 [SR:bake reading | 13.6 KD Lr:[LN:S Sacchar 34W] [AC 2r's yea g frame | PROTEIN 37909] omyces :Z28084 st] [DI | N IN MI [AC:S] cerev: 4:Y1313 | OH1-VMA5 37909] isiae] [37] ept-pln1 | INTERGENIC DB:pir2] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503001001_26757807_£2_578 | 3512 | 7284 | 951 | 316 | 1619 | 2.0e-166 |
| Description gp:[GI:e255529:g1617430] [LN:Sirepressible ABC transport system [DB:genpept-bct1] [DE:S.epiderect] [LE:1621] [RE:2550] [DI:direct] | em] [OR: midis ge | Staphyl | ococcus | s epid | ermidis] | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_26834387_f1_234 | 3513 | 7285 | 711 | 236 | 115 | 1.2e-06 |
| Description | 7 | | T-4 | | | |
| <pre>gp:[GI:g3043880] [LN:LLU95841] [OR:Lactococcus lactis] [DB:general transmembrane protein Tmp6 general signal peptide-less] [LE:<1]</pre> | npept-bo e, parti | t2] [DE al cds. | :Lactor | coccus identia | lactis | - |

sp:[LN:YVCK_BACSU] [AC:006974] [GN:YVCK] [OR:BACILLUS SUBTILIS]
[DE:HYPOTHETICAL 34.7 KD PROTEIN IN CRH-TRXB INTERGENIC REGION] [SP:006974]
[DB:swissprot] >gp:[GI:e313027:g1945651] [LN:BSZ94043] [AC:Z94043]
[PN:hypothetical protein] [GN:YVCK] [OR:Bacillus subtilis] [DB:genpept-bct1]
[DE:B.subtilis genomic DNA fragment (88 kb).] [NT:similar to hypothetical
SYCSLRA] [SP:006974] [LE:10662] [RE:11615] [DI:direct]

| SYCSLRA] [SP:006 | 974] [HE:10001] | | 7.7 | | - ***1,10 |
|-------------------|-----------------|-------|----------|----------|----------------|
| | | AA ID | NT AA LN | Score | <u>p-Value</u> |
| | NT ID | AR ID | 1111 | 1652 | 6.0e-64 |
| ORF Name | 3515 | 7287 | 014 337 | | L |
| A17503001001_2752 | 2262_c3_1426 | | | | |
| - mintion | | | | (ATP-bir | nding |

pir:[LN:G69669] [AC:G69669] [PN:choline ABC transporter (ATP-binding protein) opuBA] [GN:opuBA] [CL:glycine betaine/proline transport protein prov: ATP-binding cassette homology:CBS homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:el186061:g2635886] [LN:BSUB0018] [AC:Z99121:AL009126] [PN:choline ABC transporter (ATP-binding protein)] [GN:opuBA] [FN:high affinity transport of choline] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to [DE:Bacillus subtilis complete genome (prov)] [LE:61618] [RE:62763] [DI:complement] >gp:[GI:g2293447] [LN:AF008930] [AC:AF008930] [PN:ATPase] [GN:opuBA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis] [CB:genpept-bct2] [DE:Bacillus protein choline transport system including ATPase(opuBA), transmembrane protein (opuBA), choline binding proteinprecursor (opuBC) and transmembrane protein (opuBD) genes, completecds; and unknown gene.] [NT:OpuBA; part of choline uptake system] [LE:881] [RE:2026] [DI:direct]

| ORF Name | NT ID | | NT | 7.7 | | |
|-----------------------------|-------|--------------|-----|-----------------|-------|---------|
| AI7503001001_275268_c2_1225 | | <u>AA·ID</u> | LN | <u>AA</u> LN | Score | P-Value |
| Description | 3316 | 7288 | 125 | 374 | 766 | 5.0e-76 |

sp:[LN:HIS8_BACSU] [AC:P17731:O32008] [GN:HISC:HISH] [OR:BACILLUS SUBTILIS] [EC:2.6.1.9] [DE:PHOSPHATE TRANSAMINASE)] [SP:P17731:O32008] [DB:swissprot] >pir:[LN:A26532] [AC:A26532:G22794:H69640] [PN:histidinol-phosphate transaminase, / tyrosine and phenylalanine aminotransferase hisC:histidinol-phosphate aminotransferase] [GN:hisC:hisH] [CL:probable histidinol-phosphate transaminase] [OR:Bacillus subtilis] [EC:2.6.1.9] [DB:pir2] [MP:205 (degrees)] >gp:[GI:g143814] [LN:BACVARGNS] [AC:M80245:M15409] [PN:HisH] [GN:hisH] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:B.subtilis dbpA, mtr(A,B), gerC(1-3), ndk, cheR, aro(B,E,F,H),trp(A-F), hisH, and tyrA genes, complete cds.] [LE:14250] [RE:15332] [DI:direct] >gp:[GI:e1183707:g2634680] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:tyrosine/phenylalanine aminotransferase] [GN:hisC] [FN:histidine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.6.1.9:2.6.1.5] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:alternate gene name: aroJ; histidinol-phosphate] [SP:P17731] [LE:174116] [RE:175198] [DI:complement]

ORF Name

NT ID AA ID NT AA Score P-Value

AI7503001001_2917200_f1_277

Description

NT ID AA ID LN Score P-Value

NO-HIT

| AI7503001001 2002 | NT ID AA ID NT AA Score P-Value |
|---|---|
| 17503001001_2926425_f2_4 Description | 465 3518 7290 960 319 456 3.5e-43 |
| [OR:Bacillus subtilis] [IAC:L25604] [GN:bmrU] [FI subtilis] [DB:genpept-bc:cransporter (bmr) and its pranched-chain 2-oxo acid RE:1120] [DI:direct] >grac:D84432:D82370] [PN:Bm strain:JH642(trpC2 PheAl NA, 283 Kb region contain DI:complement] >gp:[GI:eAC:Z99116:AL009126] [PN:DR:Bacillus subtilidation] | P39074] [GN:BMRU] [OR:BACILLUS SUBTILIS] [DE:BMRU] S:swissprot] >pir:[LN:F69595] [AC:F69595] protein cotranscribed with bmr bmrU] [GN:bmrU] DB:pir2] >gp:[GI:g409286] [LN:BACBMRURBE] N:unknown, but cotranscribed with bmr] [OR:Bacillus t1] [DE:Bacillus subtilis bmrU, multidrug efflux sregulator (bmrR) genes, complete cds, and ddehydrogenase (bfmB) gene, 3' end.] [LE:227] pr[GI:d1013282:g1303947] [LN:BACJH642] mrU] [OR:Bacillus subtilis] [SR:Bacillus subtilis ning skin element.] [LE:225092] [RE:225985] 1185668:g2634834] [LN:BSUB0013] multidrug resistance protein] [GN:bmrU] B:genpept-bct1] [DE:Bacillus subtilis complete : from 2395261to 2613730.] [SP:P39074] [LE:97634] |
| E:98527] [DI:direct] | . 110m 2395261to 2613730.] [SP:P39074] [LE:97634] |
| Name 503001001_2929517_c3_139 escription -HIT | <u>NT ID AA ID <u>NT</u> <u>AA</u> <u>Score P-Value</u> 19 3519 7291 126 41</u> |
| Name 03001001_29376503_f2_504 scription HIT | NT ID AA ID NT AA Score P-Value 3520 7292 246 81 |
| Name 03001001_29378425_f3_790 cription | NT ID AA ID NT AA Score P-Value 3521 7293 132 43 |
| | |
| | |
| | |
| | |
| | |

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 A17503001001_29501510_c1_950
 3522
 7294
 1125
 374
 432
 1.2e-40

Description

sp:[LN:TAGB BACSU] [AC:P27621] [GN:TAGB] [OR:BACILLUS SUBTILIS] [DE:TEICHOIC ACID BIOSYNTHESIS PROTEIN B PRECURSOR] [SP:P27621] [DB:swissprot] >pir:[LN:C49757] [AC:C49757:C69720] [PN:polyglycerol phosphate techoic acid biosynthesis protein tagB] [GN:tagB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g143725] [LN:BACTAGABCD] [AC:M57497] [GN:tagB] [OR:Bacillus subtilis] [SR:B.subtilis (strain 168) DNA] [DB:qenpept-bct1] [DE:B.subtilis tagA, tagB, tagC and tagD genes, complete cds.] [NT:putative] [LE:1603] [RE:2748] [DI:direct] >gp:[GI:e1184482:g2636102] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:tagB] [FN:polyglycerol phosphate assembly and export] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P27621] [LE:84143] [RE:85288] [DI:direct] >qp:[GI:e1184482:q2636102] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:tagB] [FN:polyglycerol phosphate assembly and export] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P27621] [LE:84143] [RE:85288] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|-------------------------------|-------|-------|----------|-----------------|--------------|---------|
| AI7503001001_29570877_c3_1414 | 3523 | 7295 | 153 | 50 | 7 | |
| Description | | | | · • | - | |
| NO-HIT | | | | | | |
| | | - | NT | AA | | _ |
| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
| AI7503001001_3001313_c2_1262 | 3524 | 7296 | 581 | 226 | 154 | 3.6e-11 |
| Description | | | | | | |

pir:[LN:A65001] [AC:A65001] [PN:hypothetical protein b2291] [OR:Escherichia coli] [DB:pir2] >gp:[GI:g1788628] [LN:AE000318] [AC:AE000318:U00096] [PN:putative alpha helix protein] [GN:b2291] [FN:phenotype; Not classified] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 208 of 400 of the completegenome.] [NT:o199] [LE:4988] [RE:5587] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|--|---|--|-------------------------------------|--|---------------------------------|
| AI7503001001_30198587_f3_629 | 3525 | 7297 | 258 | 85 | 306 | 2.8e-27 |
| Description | | · · · · · · · · · · · · · · · · · · · | | | | |
| pir:[LN:C70024] [AC:C70024] [1]] [CL:conserved hypothetical not subtilis] [DB:pir2] >gp:[GI:el:[AC:Z99120:AL009126] [GN:yut1] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:steen [RE:112779] [DI:direct] | ifU-like 184301:g [FN:unk subtili | protei 2635719 nown] [s compl | n HP149] [LN:I OR:Bac: ete gen | 92] [O BSUB00 illus nome (| R:Bacill 17] subtilis section | lus 3] 17 of 21): |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_30265692_c3_1521 | 3526 | 7298 | 1050 | 349 | 1070 | 3.1e-108 |
| Description | | | | | | |
| gp:[GI:g4530240] [LN:AF101234] aureus] [DB:genpept-bct2] [DE:S sequence; andunknown gene.] [NT [LE:51] [RE:1010] [DI:direct] | Staphylo | coccus | aureus | dltAB | CD operc | on, complete |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503001001_30272661_f1_265 | 3527 | 7299 | 174 | 57 |] | |
| Description | | • | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_30491275_c1_961 | 3528 | 7300 | 1104 | 367 | 391 | 2.7e-36 |
| Description pir: [LN:H69867] [AC:H69867] [Figure 1.5] [IN:BSUB0008] [AC:Z99111:AL0091] [Subtilis] [DB:genpept-bct1] [DF:G6 21): from 1394791to 1603020. | is] [DB: 126] [GN E:Bacill | pir2] > [:ykvI] us subt | gp:[GI: [FN:unk ilis co | ell84: known] omplete | 961:g263 OR:Bac genome | 3742] cillus c (section 8 |
| B. subtilis] [LE:42768] [RE:438 | | | | | P | |

AΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 30601588 c3 1356 7301 834 3529 277 561 2.6e-54 Description

sp:[LN:TAGG_BACSU] [AC:P42953] [GN:TAGG] [OR:BACILLUS SUBTILIS] [DE:TEICHOIC
ACID TRANSLOCATION PERMEASE PROTEIN TAGG] [SP:P42953] [DB:swissprot]
>pir:[LN:S69202] [AC:S69202:H69720] [PN:teichoic acid permease
tagG:integral membrane protein tagG] [GN:tagG] [OR:Bacillus subtilis]
[DB:pir2] >gp:[GI:g755152] [LN:BSU13832] [AC:U13832] [PN:highly hydrophobic
integral membrane protein] [GN:tagG] [FN:teichoic acid translocation]
[OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 highly
hydrophobic integral membrane protein(tagG) gene and ATP-binding protein
(tagH) gene, complete cds.] [LE:287] [RE:1114] [DI:direct]
>gp:[GI:e1184477:g2636097] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:permease]
[GN:tagG] [FN:teichoic acid translocation] [OR:Bacillus subtilis]
[DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21):
from 3597091to 3809700.] [SP:P42953] [LE:77137] [RE:77964] [DI:complement]
>gp:[GI:e1184477:g2636097] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:permease]
[GN:tagG] [FN:teichoic acid translocation] [OR:Bacillus subtilis]

[DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 A17503001001_30661260_c1_1080
 3530
 7302
 132
 43

 Description

3597091to 3809700.] [SP:P42953] [LE:77137] [RE:77964] [DI:complement]

NO-HIT

NT AΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 30745328 c1 1103 3531 606 201 7303 200 4.8e-16

Description

pir:[LN:S56619] [AC:S56619:B65255] [PN:gpmB protein:hypothetical protein o215b] [GN:gpmB] [CL:phosphoglycerate mutase homology] [OR:Escherichia coli] [DB:pir2] >gp:[GI:g537235] [LN:ECOUW93] [AC:U14003] [OR:Escherichia coli] [DB:genpept-bct1] [DE:Escherichia coli K-12 chromosomal region from 92.8 to 00.1 minutes.] [NT:Kenn Rudd identifies as gpmB] [LE:324630] [RE:325277] [DI:direct] >gp:[GI:g1790856] [LN:AE000509] [AC:AE000509:U00096] [PN:phosphoglyceromutase 2] [GN:gpmB] [FN:enzyme; Energy metabolism, carbon: Glycolysis] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 399 of 400 of the completegenome.] [LE:8985] [RE:9632] [DI:direct]

NO-HIT

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|--|--|---|---|-----------------------------------|--|
| AI7503001001_31257943_c2_1156 | 3532 | 7304 | 414 | 137 | 669 | 9.5e-66 |
| Description | | | | | | |
| gp:[GI:g1913907] [LN:SAU91741] acid biosynthesis] [OR:Staphylo [DE:Staphylococcus aureus teich and TagX and TagD genes, completed [LE:1534] [RE:1932] [DI: | ococcus hoic aci ete cds. | aureus] .d biosy | [DB:g /nthesi | enpept s TagB | -bct1] gene, p | partialcds |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_31267503_c2_1168 | 3533 | 7305 | 813 | 270 | 784 | 6.2e-78 |
| Description | | | | | | |
| homolog ytsC] [GN:ytsC] [CL:Arsubtilis] [DB:pir2] >gp:[GI:e1:[AC:Z99119:AL009126] [GN:ytsC] [DB:genpept-bct1] [DE:Bacillus from 2997771to 3213410.] [NT:s:protein)] [LE:112638] [RE:11339 [LN:AF008220] [AC:AF008220] [PRsubtilis] [DB:genpept-bct2] [DI:LE:67028] [RE:67789] [DI:direction of the content of the | 185911:g [FN:unk subtili imilar t 99] [DI: N:transp E:Bacill | [2635522 [nown] [s comp] [o ABC to [complements] | [LN: COR:Bac Lete ge: cransponent] > [GN:yt | BSUB00 illus nome (rter (gp:[GI sC] [0 | subtilissection ATP-bind :g229317 | s] 16 of 21): ding 77] lus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_31289687_c1_1051 | 3534 | 7306 | 681 | 226 | 132 | 5.1e-06 |
| Description | | | | | | |
| <pre>gp:[GI:g4894306] [LN:AF065404] anthracis] [DB:genpept-bct2] [I complete sequence.] [LE:106772]</pre> | DE:Bacil | lus ant | hracis | virul | | |
| | | | | | | |

| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | <u>AA</u> LN | Score | P-Value |
|---|--|--|---|--|---|--|
| A17503001001_31578_c2_1265 | 3536 | 7308 | 537 | 178 | 463 | 6.4e-44 |
| Description | | | , | | | <u> </u> |
| pir:[LN:H70044] [AC:H70044] []] [OR:Bacillus subtilis] [DB:pi [AC:Z99121:AL009126] [GN:yvoF] [DB:genpept-bct1] [DE:Bacillus from 3399551to 3609060.] [NT:si [RE:191316] [DI:complement] >gp [PN:putative acetyltransferase] [DB:genpept-bct2] [DE:Bacillus [LE:36517] [RE:37035] [DI:direct | ir2] >gp [FN:unk subtili imilar t p:[GI:g2] [GN:yv subtili | o:[GI:el nown] [s compl o O-ace 618861] roF] [OR | 186184: OR:Baci ete gen tyltran [LN:AF :Bacill | g2636 llus ome (sfera 01711 us su | 009] [LN subtilis section se] [LE: 3] [AC:A btilis] | EBSUB0018]] 18 of 21): 190798] F017113] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_31803377_£3_884 | 3537 | 7309 | 612 | 203 |] | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_31876563_c1_1124 | 3538 | 7310 | 1239 | 412 | 1924 | 9.7e-199 |
| Description | | • | | | | |
| gp:[GI:d1013747:g1405336] [LN:I transporter] [GN:dltB] [OR:Stap (isolate:KAN96) DNA] [DB:genper unkown function and dlt operon [NT:Sequence homologous to the [DI:direct] >gp:[GI:g4530242] membrane protein DltB] [GN:dltB teichoic] [OR:Staphylococcus au aureus dltABCD operon, complete [RE:4258] [DI:direct] | phylococ pt-bct1] dltA,dl dltB ge [LN:AF10 3] [FN:i ireus] [| Cus aur [DE:St tB, dlt nes of] 1234] [nvolved DB:genp | eus] [S aphyloc C and d [LE:40 AC:AF10 in D-a ept-bct | R:Staj occus ltD ge 71] [1 1234] lanine 2] [D] | phylococ aureus enes,com RE:5285] [PN:put e transf E:Staphy | cus aureus gene for plete cds.] ative er into lococcus |
| ORF Name | NT ID | AA ID | NT | AA | Score | P-Value |
| A17503001001 32062553 c1 943 | 3539 | 7311 | <u>LN</u> 468 | <u>LN</u> 155 | 354 | 2.3e-32 |
| Description | | | | | | |
| pir:[LN:S61393] [AC:S61393] [R [SR:strain C-125, , strain C-12 >gp:[GI:d1007182:g854656] [LN:R sp.] [SR:Bacillus sp. (strain:C Na+/H+ antiporter system responsible gene] [LE:2669] [RR | 25] [SR: BACAPS] C-125) D nsible g | strain [AC:D31 NA] [DB enes.] | C-125, 823] [P: :genpep [NT:Na+ |] [DB N:ORF2 t-bct] | :pir2] 2] [OR:B 1] [DE:B | acillus acillus sp. |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|--|--|---|--|---|--|
| AI7503001001_32070827_£3_789 | 3540 | 7312 | 168 | 55 | 7 | |
| Description | | , | | | | |
| NO-HIT | | | | | | |
| | | | NT | AA | | |
| ORF Name | NT ID | AA ID | LN | LN | Score | <u>P-Value</u> |
| AT7503001001_32212902_c2_1135 | 3541 | 7313 | 1035 | 344 | 468 | 1.9e-44 |
| Description | | | | | | |
| [AC:AJ223978] [PN:putative hemisubtilis] [DB:genpept-bct1] [DB:yvsA to yvqA.] [LE:24699] [RE:2 [LN:BSUB0017] [AC:Z99120:AL0091 subtilis] [DB:genpept-bct1] [DB:17 of 21): from 3197001to 34144 [LE:204527] [RE:205588] [DI:com [LN:BSUB0018] [AC:Z99121:AL0091 subtilis] [DB:genpept-bct1] [DB:genpept-bct1] [DB: of 21): from 3399551to 36090 [RE:3038] [DI:complement] | E:Bacill 25760] [126] [GN E:Bacill 120.] [N mplement 126] [GN E:Bacill | us subt DI:dire [:yvrB] us subt T:simil] >gp:[[:yvrB] us subt | ilis 42 ct] >gp [FN:unk ilis co ar to i GI:e118 [FN:unk ilis co | 2.7kB 1 c:[GI:cnown] cmpletcron possible from possible fro | DNA fragel 184396 [OR:Bacel genome grmease] g2635830 [OR:Bacel genome | gment from 5:g2635814] cillus c (section cillus cillus cillus c (section |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_32225012_f1_54 | 3542 | 7314 | 498 | 165 | 276 | 4.2e-24 |
| Description pir: [LN:F69870] [AC:F69870] [EGN:ykzA] [CL:hypothetical property of the property | tein yk :BSAJ257 :lis] [D xlyA an RE:34555 :BSUB000 :lis] [D 21): fro | lA] [OR 1] [AC: B:genpe d ykoR.] [DI:d 7] [AC: B:genpe m 11943 | :Bacill AJ00257 pt-bct1] [NT:h irect] Z99110: pt-bct1 91to 14 | .us sul /1] [PI .] [DE nomolos AL009: .] [DE | otilis] N:YknA] :Bacillu gous to 126] [GN :Bacillu .] [NT:a | [DB:pir2] [GN:yknA] us subtilis OsmC from U:ykzA] us subtilis ulternate |
| ORF Name AI7503001001_32475037_f3_651 Description | NT ID | <u>AA ID</u> 7315 | NT LN 123 | AA LN 40 | Score | <u>P-Value</u> |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
|---|--|---|--|---|--|--|
| A17503001001_32611068_c1_927 | 3544 | 7316 | 1674 | 557 | 1694 | 2.3e-174 |
| Description | | | | | | |
| sp:[LN:SYR_BACSU] [AC:P46906] [DE:ARGINYL-TRNA SYNTHETASE, ([DB:swissprot] >pir:[LN:E69589] ligase, argS:arginyl-tRNA synt ligase] [OR:Bacillus subtilis] >gp:[GI:e1186234:g2636270] [LN [PN:arginyl-tRNA synthetase] [[DB:genpept-bct1] [EC:6.1.1.19] (section 20 of 21): from 37984 [RE:35973] [DI:complement] >gp [AC:Z97024] [PN:arginyl tRNA some second | ARGININE ARG | ETRNA 59589:S6 [GN:arg .1.19] 00] [AC: [OR:Ba acillus .0550.] 4388:g2 se] [GN: .s ywiA, | LIGASE) 0082] S] [CI [DB:pin Z99123: cillus subtili [SP:P46 224756] argS] sbo, y | (ARGI [PN:ar L:Baci 2] AL009: subtil is comp [S906] [LN:I | RS)] [SPrginine- llus arg 126] lis] plete ge [LE:3430 3SZ97024 | :P46906] -tRNA ininetRNA nome 3] j ubtilis] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_33210952_£2_370 | 3545 | 7317 | 234 | 77 | 93 | 0.0025 |
| Description | | | | | | |
| <pre>gp:[GI:g3859891] [LN:AF072678] [OR:Trichomonas vaginalis] [DB alpha-actinin (AACTI) mRNA, con [LE:247] [RE:3042] [DI:direct]</pre> | :genpept | -inv2] | [DE:Tri | chomor | nas vagi | nalis |
| ORF Name AI7503001001_33239001_f2_356 | NT ID | AA ID | NT LN 276 | <u>AA</u> <u>LN</u> 91 | Score | P-Value |
| Description | | | | | | |
| NO-HIT | . <u>-</u> | | | | | |
| ORF Name AI7503001001_33304063_f1_151 Description | NT ID | <u>AA ID</u> 7319 | <u>NT</u> <u>LN</u> 159 | <u>AA</u> <u>LN</u> 52 | <u>Score</u> | P-Value |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|---|---|--|--|--|-------------|
| A17503001001_33414143_c3_1395 | 3548 | 7320 | 1710 | 569 | 619 | 1.9e-60 |
| Description | | | | | | |
| <pre>gp:[GI:g2773332] [LN:AF040718] [GN:cydC] [OR:Shigella flexner transporter CydC (cydC) gene, [LE:1648] [RE:3369] [DI:direct</pre> | i] [DB:g | genpept-l | bct2] | [DE:Sh | igella f | lexneri ABC |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_33628441_f3_645 | 3549 | 7321 | 255 | 84 | | |
| Description | | | | | | |
| NO-HIT | | | _ | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_33645967_f1_18 | 3550 | 7322 | 282 | 93 | 73 | 0.019 |
| Description | | | | | | |
| | | | | | | |
| <pre>gp:[GI:g2444074] [LN:CPU88070] of type III secretory] [OR:Chl [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini</pre> | amydophi cds2, c | la cavia opN and | ae] [Di scc1 q | B:genp genes, | ept-bct2 complet | 2] |
| of type III secretory] [OR:Chl [DE:Chlamydophila caviae cds1, | amydophi cds2, c | la cavia opN and | ae] [Di scc1 q | B:genp genes, | ept-bct2 complet | 2] |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. | amydophi cds2, c a] [LE:4 | la cavia copN and 766] [RI AA ID | ae] [Di scc1 g E:5206] | B:genp genes,] [DI: | ept-bct2 complet direct] | e cds.] |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name | amydophi cds2, c a] [LE:4 NT ID | la cavia copN and 766] [RI AA ID | ae] [Di scc1 g E:5206] <u>NT</u> <u>LN</u> | B:genp genes,] [DI: <u>AA</u> <u>LN</u> | ept-bct2 complet direct] | e cds.] |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersinian Corporation of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersinian of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [OR:Chl. [DE:Chl. [DE: | amydophi cds2, c a] [LE:4 NT ID | la cavia copN and 766] [RI AA ID | ae] [Di scc1 g E:5206] <u>NT</u> <u>LN</u> | B:genp genes,] [DI: <u>AA</u> <u>LN</u> | ept-bct2 complet direct] | e cds.] |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name AI7503001001_34069680_c1_1125 Description | amydophi cds2, c a] [LE:4 NT ID | la cavia copN and 766] [RI AA ID | ae] [Di scc1 g E:5206] <u>NT</u> <u>LN</u> | B:genpgenes, CDI: AA LN 70 | ept-bct2 complet direct] | e cds.] |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name A17503001001_34069680_c1_1125 Description NO-HIT | amydophi cds2, c a] [LE:4 NT ID | la cavia copN and 766] [RI AA ID 7323 | nae] [Di scc1 9 E:5206 NT LN 213 | B:genp genes,] [DI: <u>AA</u> <u>LN</u> | ept-bct2 complet direct] Score | P-Value |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersinis ORF Name AI7503001001_34069680_c1_1125 Description NO-HIT ORF Name | amydophicds2, cal [LE:4 | la cavia copN and 766] [RI AA ID 7323 | ne] [Di scc1 scc1 scc1 scc1 scc1 scc1 scc1 scc | B:genpgenes, [DI: AA LN 70 AA LN | ept-bct2 complet direct] Score | P-Value |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name AI7503001001_34069680_c1_1125 Description NO-HIT ORF Name AI7503001001_34160625_c3_1368 | amydophicds2, cal [LE:4 | la cavia copN and 766] [RI AA ID 7323 | ne] [Di scc1 scc1 scc1 scc1 scc1 scc1 scc1 scc | B:genpgenes, [DI: AA LN 70 AA LN | ept-bct2 complet direct] Score | P-Value |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name AI7503001001_34069680_c1_1125 Description NO-HIT ORF Name AI7503001001_34160625_c3_1368 Description | amydophicds2, cal [LE:4 | la cavia copN and 766] [RI AA ID 7323 | ne] [Di scc1 scc1 scc1 scc1 scc1 scc1 scc1 scc | B:genpgenes, [DI: AA LN 70 AA LN | ept-bct2 complet direct] Score | P-Value |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name AI7503001001_34069680_c1_1125 Description NO-HIT ORF Name AI7503001001_34160625_c3_1368 Description NO-HIT | amydophicds2, cal [LE:4 NT ID 3551 NT ID 3552 | AA ID 7324 AA ID | NT LN 438 | B:genpgenes, CDI: AA LN 70 AA LN 145 | ept-bct2 complet direct] Score Score | P-Value |
| of type III secretory] [OR:Chl. [DE:Chlamydophila caviae cds1, [NT:similar to SycE of Yersini. ORF Name AI7503001001_34069680_c1_1125 Description NO-HIT ORF Name AI7503001001_34160625_c3_1368 Description NO-HIT ORF Name | amydophicds2, cal [LE:4 NT ID 3551 NT ID 3552 | AA ID 7324 AA ID | NT LN 438 | B:genpgenes, DI: AA LN 70 AA LN 145 | ept-bct2 complet direct] Score Score | P-Value |

| ORF Name | NT ID | AA ID | LN | <u>AA</u> LN | Score | P-Value |
|---|--|--|--|--|--|------------------------------|
| A17503001001_34173385_c1_975 | 3554 | 7326 | 621 | 206 | 151 | 7.4e-11 |
| Description | | | | | | |
| pir:[LN:E71040] [AC:E71040] [[OR:Pyrococcus horikoshii] [DB [LN:AP000006] [AC:AP000006:AB005215:AB009510 [PN:166aa long hypothetical pr [SR:Pyrococcus horikoshii (str [DB:genpept-bct1] [DE:Pyrococcus horikoshii)] | :pir2] > :AB00951 otein] [ain:OT3) us horik | gp:[GI: 1:AB009 GN:PH16 DNA, coshii O | d103160 512:AB0 13] [OI lone:Py | 68:g32 009513 R:Pyro yrococ omic D | :AB00951 coccus h cus hori NA, 1166 | .4] orikoshii] .koshi] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_34173750_c3_1485 | 3555 | 7327 | 480 | 159 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_34176550_f1_24 | 3556 | 7328 | 132 | 43 | | |
| Description NO-HIT | | | | | | |
| | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
| AI7503001001_34189817_c1_945 | 3557 | 7329 | 2052 | 683 | 888 | 5.9e-89 |
| Description pir:[LN:E70040] [AC:E70040] [PN:conserved hypothetical protein yvgP] [GN:yvgP] [CL:hypothetical protein yvgP] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1186030:g2635855] [LN:BSUB0018] [AC:Z99121:AL009126] [GN:yvgP] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:similar to hypothetical proteins] [LE:27843] [RE:29855] [DI:complement] | | | | | | |

| ORF Name | NT ID | AA ID | LN | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|--|--|--|---------------------------------|---|---|
| A17503001001_34242162_c3_1511 | 3558 | 7330 | 489 | 162 | 529 | 6.5e-51 |
| Description | | | | | | |
| <pre>pir:[LN:E70019] [AC:E70019] [] [CL:Yeast nitrogen fixation [OR:Bacillus subtilis] [DB:pir [AC:Z99120:AL009126] [GN:yurV] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:s [RE:159513] [DI:complement]</pre> | protein: 2] >gp:[[FN:unk subtili | nitroge GI:e118 nown] [s compl | n fixat 4346:g2 OR:Baci ete gen | ion p 63576 llus ome (| rotein h 4] [LN:B subtilis section | omology] SUB0017]] 17 of 21): |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_34251887_c2_1322 | 3559 | 7331 | 1266 | 421 | 1649 | 1.3e-169 |
| Description gp:[GI:g4530244] [LN:AF101234] [AC:AF101234] [PN:putative exoprotein DltD] [GN:dltD] [FN:involved in D-alanine transfer into teichoic] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus dltABCD operon, complete sequence; andunknown gene.] [LE:4509] [RE:5684] | | | | | | |
| ORF Name A17503001001_34254581_c1_954 Description NO-HIT | NT ID | AA ID | NT LN 147 | <u>AA</u> <u>LN</u> 48 | Score | P-Value |
| | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | $\frac{AA}{LN}$ | Score | P-Value |
| <u>Description</u> | 3561 | 7333 | 312 | 103 | 75 | 0.018 |
| <pre>gp:[GI:g488925] [LN:A13473] [Additional content of the conten</pre> | te P. fa | lciparu | m] [DB: | genpe | pt-pat] | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|-------------------------------------|---|--|--------------------------------------|---|---|
| AI7503001001_34562762_c3_1336 | 3562 | 7334 | 750 | 249 | 230 | 3.2e-19 |
| Description | | | | | | |
| pir:[LN:D71157] [AC:D71157] [F [OR:Pyrococcus horikoshii] [DB: [LN:AP000002] [AC:AP000002:AB009475:AB009476: [PN:232aa long hypothetical pro [SR:Pyrococcus horikoshii (stra horikoshii OT3 genomic DNA, 287 [RE:133673] [DI:direct] | :pir2] > :AB00947 otein] [ain:OT3) | gp:[GI: 7:AB009 [GN:PH04 DNA] [| d103048 478:AB(59] [OI DB:genj | 38:g32 009479 R:Pyro pept-b | 56862] :AB00948 coccus h ct1] [DE | 80] norikoshii] E:Pyrococcus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_34571000_f1_163 | 3563 | 7335 | 537 | 178 | 393 | 1.7e-35 |
| Description sp:[LN:ADH2_ENTHI] [AC:Q24803:Q [EC:1.1.1.1:1.2.1.10] [DE:DEHYI [DB:swissprot] >gp:[GI:g488430] dehydrogenase 2] [OR:Entamoeba histolytica HM1:IMSS alcohol de [NT:The derived amino acid sequ [DI:direct] | DROGENAS LIN:EH histoly hydroge | E, (ACD [U04863] rtica] [nase 2 | H)] [SI [AC:U(DB:genr (EhADH2 | 9:Q248 04863] pept-i 2)mRNA | 03:Q2764 [PN:alc nv1] [DE , comple | 9] cohol E:Entamoeba ete cds.] |
| ORF Name AI7503001001_34589027_c2_1197 Description | NT ID | <u>AA ID</u> 7336 | <u>NT</u> <u>LN</u> 315 | <u>AA</u> <u>LN</u> 104 | Score | P-Value 0.00023 |
| sp:[LN:Y420_METJA] [AC:Q57863] [DE:HYPOTHETICAL PROTEIN MJ0420 [AC:D64352] [PN:hypothetical r [DB:pir2] [MP:FOR378394-379536 [AC:U67494:L77117] [PN:O-antige |] [SP:Q protein] >gp:[| 57863] MJ0420] GI:g159 | [DB:swi [OR:Me 1123] | sspro thano LN:U6 | t] >pir: coccus j 7494] | [LN:D64352] |

[OR:Methanococcus jannaschii] [DB:genpept-bct2] [DE:Methanococcus jannaschii

section 36 of 150 of the complete genome.] [NT:similar to GB:M60066

SP:P26479 PID:154343 percent] [LE:1855] [RE:2997] [DI:direct]

NTAA ORF Name NT ID AA ID Score P-Value LN LN AI7503001001_34612887_f2_512 465 154 3565 7337 260 2.1e-22

Description

pir: [LN:E69857] [AC:E69857] [PN:conserved hypothetical protein ykmA] [GN:ykmA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1181515:g2632035] [LN:BSAJ2571] [AC:AJ002571] [PN:YkmA] [GN:ykmA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 56 kb DNA fragment between xlyA and ykoR.] [LE:33559] [RE:34002] [DI:complement] >qp:[GI:e1183335:q2633669] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:ykmA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [NT:similar to hypothetical proteins] [LE:186508] [RE:186951] [DI:complement]

NT AΑ ORF Name NT ID Score AA ID P-Value LN LN AI7503001001 34627136 f3 681 3566 7338 306 101 73 0.014

Description

pir:[LN:S36690] [AC:S36690] [PN:hypothetical protein] [OR:Autographa californica nuclear polyhedrosis virus:AcMNPV] [DB:pir2] >gp:[GI:g296321] [LN:ACNPVDNA] [AC:X71415] [GN:ORF 339] [OR:Autographa californica nucleopolyhedrovirus] [DB:genpept-vrl] [DE:Autographa californica nuclear polyhedrosis virus DNA.] [SP:Q06669] [LE:<1] [RE:339] [DI:complement]

NT AA ORF Name NT ID AA ID Score P-Value LN LN 312 AI7503001001 34641875 c3 1406 939 3567 7339 1.9e-69 704

Description

pir: [LN:A69627] [AC:A69627] [PN:fructose 1-phosphate kinase fruB] [GN:fruB [CL:6-phosphofructokinase 2] [OR:Bacillus subtilis] [DB:pir2] >qp: [GI:e1185029:q2633810] [LN:BSUB0008] [AC:Z99111:AL009126] [PN:fructose-1-phosphate kinase] [GN:fruB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.56] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:112945] [RE:113856] [DI:direct] >gp:[GI:g3282124] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:fructose-1-phosphate kinase] [GN:fruB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to L. lactis tagatose-6-phosphate kinase,] [LE:13433] [RE:14344] [DI:direct]

| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | AA LN | Score | P-Value |
|--|---|--|---|-------------------------------------|---------------------------------------|----------------------------|
| AI7503001001_34642567_c1_1089 | 3568 | 7340 | 2406 | 801 | 2175 | 1.0e-228 |
| Description | · | | | | | |
| pir:[LN:G70027] [AC:G70027] [GN:yvaJ] [CL:virulence-assoce subtilis] [DB:pir2] >gp:[GI:e1 [AC:Z99121:AL009126] [GN:yvaJ] [DB:genpept-bct1] [DE:Bacillus from 3399551to 3609060.] [NT:series [RE:53714] [DI:complement] | iated pr 186049:9 FN:unk subtili | rotein v 12635874 (nown] Ls compl | racB hor [LN:] [OR:Bac: .ete ger | molog] BSUB00 illus nome (| [OR:Bac 18] subtilis section | cillus s] 18 of 21): |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_34645311_f3_867 Description | 3569 | 7341 | 525 | 174 | 186 | 1.4e-14 |
| <pre>gp:[GI:g4981094] [LN:AE001732] hypothetical protein] [GN:TM05 [DE:Thermotoga maritima section [NT:similar to SP:P46854 PID:6] [DI:complement]</pre> | 77] [OR: on 44 of | Thermot | oga mar | ritima mplete |] [DB:ge | enpept-bct2] .] |
| ORF Name AI7503001001_3518_f2_375 Description NO-HIT | NT ID | <u>AA ID</u> 7342 | <u>NT</u> <u>LN</u> 159 | <u>AA</u> <u>LN</u> 52 | <u>Score</u> | P-Value |
| | | | | | | |
| ORF Name | NT ID | AA ID | <u>NT</u> <u>LN</u> | AA LN | Score | P-Value |
| A17503001001_35422880_c2_1228 | 3571 | 7343 | 129 | 42 | | |
| <u>Description</u> | | | | | | |
| NO-HIT | | | | | | |

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 A17503001001_35444127_f2_443
 3572
 7344
 663
 220
 514
 2.5e-49

Description

sp:[LN:YVYE BACSU] [AC:P32437:P96500] [GN:YVYE:YVHK] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 24.8 KD PROTEIN IN DEGS-TAGO INTERGENIC REGION] [SP:P32437:P96500] [DB:swissprot] >pir:[LN:A70049] [AC:A70049:A30191] [PN:conserved hypothetical protein yvyE] [GN:yvyE] [CL:hypothetical protein HI0722] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184457:g2636077] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:yvyE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: yvhK; similar to hypothetical] [SP:P32437] [LE:48723] [RE:49376] [DI:direct] >gp:[GI:g1762328] [LN:BSU56901] [AC:U56901] [PN:Ycr59c/YigZ homolog] [GN:yvhK] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis putative transcriptional regulator (yvhJ), Ycr59c/YiqZ homolog (yvhK), histidine kinase (degS), transcriptional regulator of degradation enzyme (degU), (degV), (comFA), (comFB), (comFC), flagellar protein (yviB), negative regulator of flagellin(flgM), flagellar protein (yviC), flagellar-hook associated protein1 (flgK), flagellar-hook associated protein 3 (flgL), (yviE), transmembrane protein (yviF), (csrA), flagellin (hag), flagellarprotein (yviH), flagellar hook-associated protein 2 (fliD), flagellar protein (fliS), flagellar protein (fliT), sigma-54modulator homolog (yviI), and (secA) genes, complete cds.] [LE:1217] [RE:1870] [DI:complement] >gp:[GI:e1184457:g2636077] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:yvyE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: yvhK; similar to hypothetical] [SP:P32437] [LE:48723] [RE:49376] [DI:direct]

NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 35945277 c3 1463 7345 1065 354 3573 5.8e-130 1275 Description

sp:[LN:RF2 BACSU] [AC:P28367:O34444] [GN:PRFB] [OR:BACILLUS SUBTILIS] [DE:PEPTIDE CHAIN RELEASE FACTOR 2 (RF-2)] [SP:P28367:034444] [DB:swissprot] >pir:[LN:JN0146] [AC:H69681:JN0146] [PN:translation releasing factor RF-2:peptide chain release factor 2:prfB] [GN:prfB] [CL:translation releasing factor] [OR:Bacillus subtilis] [DB:pir1] >gp:[GI:e1184435:g2636055] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:peptide chain release factor 2] [GN:prfB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P28367] [LE:29109:30139] [RE:30137:30210] [DI:complementJoin] >gp:[GI:g2331287] [LN:AF013188] [AC:AF013188] [PN:release factor 2] [GN:prfB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis release factor 2 (prfB) gene, complete cds.] [NT:orf3; naturally occurring frame-shift] [LE:75:148] [RE:146:1176] [DI:directJoin] >gp:[GI:g2618874] [LN:AF017113] [AC:AF017113] [PN:putative peptide chain release factor RF-2] [GN:prfB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis 300-304 degree genomic sequence.] [LE:83:156] [RE:154:1184] [DI:directJoin] >gp:[GI:e1184435:g2636055] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:peptide chain release factor 2] [GN:prfB] [OR:Bacillus subtilis] [DB:qenpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P28367] [LE:29109:30139] [RE:30137:30210] [DI:complementJoin]

NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001_35988961 f2 376 3574 7346 129 42 Description NO-HIT NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 36131311 c3 1455 231 3575 7347 696 194 2.1e-15

Description

gp:[GI:g451867] [LN:BACFLGMK] [AC:L14437] [OR:Bacillus subtilis]
[SR:Bacillus subtilis (strain W168) (library: lambda gtWES library]
[DB:genpept-bct1] [DE:Bacillus subtilis (clones pDM116 and pDM113) flagellin synthesisregulatory protein (flgM) and flagellar hook-filament junctionprotein (flgK) genes and orf139, orf160, orfX, complete cds's.]
[NT:in Z18629, ORFX is called comForf3; ORFX; putative] [LE:<1] [RE:754]
[DI:direct]</pre>

[RE:13436] [DI:direct]

NT ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 36133385 c3 1377 3576 7348 681 226 2.3e-16 203 Description pir:[LN:G69798] [AC:G69798] [PN:hypothetical protein yetJ] [GN:yetJ] [CL:hypothetical protein yetJ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182699:g2633033] [LN:BSUB0004] [AC:Z99107:AL009126] [GN:yetJ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [LE:186800] [RE:187444] [DI:direct] NT AΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001_36229676_c1_936 3577 264 7349 795 238 4.5e-20 Description gp:[GI:g3172115] [LN:ACCPCAOP] [AC:L05770:U04359:M33798:U20284:U11554:L13114:L03407] [PN:beta-ketoadipate enol-lactone hydrolase] [GN:pcaD] [OR:Acinetobacter sp. ADP1] [DB:genpept-bct2] [EC:3.1.1.24] [DE:Acinetobacter sp. ADP1 pca-qui-pob supraoperonic cluster, completesequence.] [NT:ELH] [LE:6843] [RE:7643] [DI:direct] NT AΑ ORF Name NT ID AA ID P-Value Score LN LN 258 A17503001001_36230252_c1_1000 3578 7350 777 480 1.0e-45 Description pir:[LN:B69627] [AC:B69627] [PN:transcription repressor of fructose operon fruR] [GN:fruR] [CL:regulatory protein gutR] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185028:g2633809] [LN:BSUB0008] [AC:Z99111:AL009126] [PN:transcriptional regulator (DeoR family)] [GN:fruR] [FN:negative regulation of the fructose operon] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:112193] [RE:112948] [DI:direct] >gp:[GI:g3282123] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:FruR] [GN:fruR] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to L. lactis lactose PTS system repressor,] [LE:12681]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|--|--|---|--|---|-------------------------------------|
| A17503001001_36367302_c3_1330 | 3579 | 7351 | 1564 | 187 | 122 | 2.7e-06 |
| Description | | JL | J L | J L | | |
| sp:[LN:Y359_METJA] [AC:Q57805] [DE:HYPOTHETICAL PROTEIN MJ035 [AC:G64344] [PN:hypothetical [DB:pir2] [MP:REV327449-326805 [AC:U67489:L77117] [PN:M. jann [GN:MJ0359] [OR:Methanococcus [DE:Methanococcus jannaschii s [NT:hypothetical protein; iden [DI:complement] | 9] [SP:Q protein] >gp: aschii p jannasch ection 3 | 257805] MJ0359] [GI:g159 predicte hii] [DB 61 of 15 | [DB:sw. [OR:Ma 1068] d codin :genpe 0 of t | isspro ethano [LN:U6 ng reg pt-bct he com | t] >pir: coccus j 7489] ion MJ03 2] plete ge | [LN:G64344] annaschii] 559] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_36563_c2_1302 | 3580 | 7352 | 1074 | 357 | 908 | 4.5e-91 |
| Description | | • | | | | - |
| homolog yusC] [GN:yusC] [CL:A subtilis] [DB:pir2] >gp:[GI:e1 [AC:Z99120:AL009126] [GN:yusC] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:s protein)] [LE:165325] [RE:1663 | 184353:g [FN:unk subtili imilar t | [2635771 [nown] [[s compl [o ABC t |] [LN:] OR:Bac: ete ger ranspo: | BSUB00 illus nome (| 17] subtilis section | :] 17 of 21): |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_36601562_c3_1403 | 3581 | 7353 | 1182 | 393 | 1669 | 1.0e-171 |
| Description sp:[LN:NORA_STAAU] [AC:P21191] [DE:QUINOLONE RESISTANCE NORA >pir:[LN:A37838] [AC:A37838] [DB:pir2] >gp:[GI:d1014850:g21:norA] [GN:norA] [OR:Staphylocottk2566) genomic DNA] [DB:genperore:[RE:1644] [DI:direct] | PROTEIN] [PN:norA 6975] [I ccus aur | [SP:P2 protei N:STANO eus] [S | 1191] n] [OR RA] [A(R:Stap) | [DB:sw :Staph C:D901 hyloco | issprot] ylococcu 19] [PN: ccus aur | s aureus] ORF for eus (strain |
| ORF Name A17503001001_37927_f1_282 Description | NT ID | <u>AA ID</u> 7354 | <u>NT</u> <u>LN</u> | AA LN 40 | Score | P-Value |

| ORF Name | NT ID | AA ID | NT | <u>AA</u> | Score | P-Value |
|---|---|--|--|--|--|--|
| A17503001001 391527_f3 880 | 3583 | 17355 | <u>LN</u> | <u>LN</u> | 700 | 4.9e-69 |
| Description | | 1,333 | 1330 | 309 | | 4.96-09 |
| sp:[LN:TAGH_BACSU] [AC:P42954] ACID TRANSLOCATION ATP-BINDING >pir:[LN:S69203] [AC:S69203:A6:ATP-binding protein tagH] [GN:COR:Bacillus subtilis] [DB:pir:[DN:ATP-binding protein] [GN:tagh] [OR:Bacillus subtilis] [DB:gen] hydrophobic integral membrane polymerates (tagh) gene, complete cds.] [LN:Sqp:[GI:e1184476:g2636096] [LN:COR:Bacillus subtilis] [DB:gen] genome (section 19 of 21): from [RE:77117] [DI:complement] >qp [AC:Z99122:AL009126] [PN:ATP-binding protein] [CR:Tanslocation] [OR:Bacillus subtilis] [DB:gen] translocation] [OR:Bacillus subtilis] [DB:Gen] [AC:Z99122:AL009126] [PN:ATP-binding protein] >qp [AC:Z99122:AL009126] [PN:ATP-binding protein] [CR:Tanslocation] [OR:Bacillus subtilis] [DB:Gen] [AC:Z99122:AL009126] [PN:ATP-binding protein] [CR:Tanslocation] [OR:Bacillus subtilis] [DB:Gen] | PROTEIN 9721] tagH] 2] >gp: agH] [FN pept-bct protein E:1134] :BSUB001 agH] [FN pept-bct m 359709 :[GI:ell inding p btilis] 21): fr | N TAGH] [PN:teic] [CL:ATP- [GI:g755] I:teicho [1] [DE: [tagG] g [RE:271 [9] [AC: I:teicho [1] [DE: Plto 380 [84476:g [DB:gen | [SP:P42] hoic ac binding 153] [1 ic acic Bacilla ene and 7] [DI Z99122 ic acic Bacilla 9700.] 2636096 [GN:ta pept] | 2954] cid tra g casse LN:BSUI d trans us subs d ATP-1 :direct :AL009 d trans us subs [SP:P4 6] [LN agH] [1 | [DB:swise anslocate to the control of the control o | ssprot] cion mology] [AC:U13832] on] 68 highly protein on] omplete [LE:75534] 19] noic acid subtilis |
| ORF Name AI7503001001_3922550_f1_67 Description NO-HIT | NT ID | <u>AA ID</u> 7356 | <u>NT</u> <u>LN</u> 138 | <u>AA</u> <u>LN</u> 45 | Score | <u>P-Value</u> |
| ORF Name | NT ID | AA ID | NT LN | AA LN | <u>Score</u> | P-Value |
| AI7503001001_3928762_f2_369 Description | 3585 | 7357 | 1368 | 455 | 839 | 9.2e-84 |
| pir:[LN:B70007] [AC:B70007] [IGN:yubG] [CL:Na+-ATP synthase] >gp:[GI:e1185983:g2635594] [LN [FN:unknown] [OR:Bacillus subtromplete genome (section 16 of Na+-transporting ATP synthase] | e chain :BSUB001 ilis] [I 21): fr | J] [OR: .6] [AC: .8:genpe .com 2997 | Bacillo Z99119: pt-bct1 771to 3 | us subt :AL0091 1] [DE: 3213410 | cilis] L26] [GN :Bacillu D.] [NT: | [DB:pir2] J:yubG] us subtilis similar to |
| ORF Name AI7503001001_3939218_c1_922 Description | NT ID | <u>AA ID</u> 7358 | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> | Score | <u>P-Value</u> |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|--|---|---|--------------------------------------|---|-------------------------------------|
| AI7503001001_3942263_c3_1381 | 3587 | 7359 | 726 | 241 | 913 | 1.3e-91 |
| Description | | | | | | |
| pir:[LN:C69793] [AC:C69793] [3 [GN:yeeI] [CL:hypothetical proposed | otein MG :BSUB000 ilis] [I 21): fro | 3332] [0: 04] [AC: 0B:genpe om 60070 | R:Bacil Z99107 pt-bctl 1 to813 | llus s :AL009 1] [DE 3890.] | ubtilis] 126] [GN :Bacillu [NT:sim | [DB:pir2] N:yeeI] us subtilis |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_3944001_c1_1130 | 3588 | 7360 | 1524 | 507 | 684 | 2.4e-67 |
| <pre>>gp:[GI:e1184284:g2635702] [LN [FN:unknown] [OR:Bacillus subt: complete genome (section 17 of leucyl aminopeptidase] [LE:984' ORF Name</pre> | ilis] [I 21): fr | B:genpercom 3197 | pt-bct1 001to 3 | L] [DE 341442 | :Bacillu 0.] [NT: | ıs subtilis |
| | | | LN | <u>LN</u> | | |
| A17503001001_3948587_f2_340 Description | 3589 | 7361 | 873 | 290 | 378 | 6.5e-35 |
| gp:[GI:g4980658] [LN:AE001701] hypothetical protein] [GN:TM016 [DE:Thermotoga maritima section [NT:similar to GB:AE000666 pero | 64] [OR: n 13 of | Thermoto 136 of | oga man | citima mplete |] [DB:ge genome. | enpept-bct2] |
| ORF Name A17503001001 3953452 c2 1201 | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> 168 | Score | P-Value 1.3e-29 |
| Description | <u> </u> | | | | J | |
| pir:[LN:D69849] [AC:D69849] [I | | _ | _ | | _ | |

[DB:pir2] >gp:[GI:e1183226:g2633560] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:yjdI] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus

[NT:similar to transcription regulation] [LE:82766] [RE:83245] [DI:direct]

subtilis complete genome (section 7 of 21): from 1194391to 1411140.]

| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | AA LN | Score | P-Value |
|--|--|---|--|--|---|--|
| A17503001001_3955067_c2_1278 | 3591 | 7363 | 126 | <u>==:</u> 41 | ٦ | |
| Description | | JL | | | _ | |
| NO-HIT | | | <u>. </u> | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AT7503001001_3959377_f3_886 | 3592 | 7364 | 126 | 41 | 149 | 1.7e-10 |
| Description | | | | | | |
| <pre>gp:[GI:e255528:g1617429] [LN:S] [FN:iron repressible ABC trans] [DB:genpept-bct1] [DE:S.epider] [LE:878] [RE:1624] [DI:direct]</pre> | port sys | stem] [OF | R:Stap | hyloco | ccus ep: | idermidis] |
| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | <u>P-Value</u> |
| | 3593 | 7365 | <u></u> | 208 | 139 | 1.1e-08 |
| A17503001001_3962915_f2_507 | | | | | 11 | |
| Description pir: [LN:C71375] [AC:C71375] [| | rved hyp | oothet | ical i | | |
| Description | OR:Trepo >gp:[GI served h idum] [I e genome | erved hyponema pal eg332228 egpotheti eggenper | pothet llidum 38] [L ical i pt-bct simila | ical i subsp N:AE00 ntegra 2] [DE r to P | . pallio 1188] l membra :Trepone | dum] [SR:, ane] ema pallidum |
| Description pir: [LN:C71375] [AC:C71375] [protein TP0033] [GN:TP0033] [GS:C71375] [syphilis spirochete] [DB:pir2] [AC:AE001188:AE000520] [PN:content of the complete of the | OR:Trepo >gp:[GI served h idum] [I e genome | erved hyponema pal eg332228 egpotheti eggenper | pothet llidum 38] [L ical i pt-bct simila | ical i subsp N:AE00 ntegra 2] [DE r to P | . pallio 1188] l membra :Trepone | dum] [SR:, ane] ema pallidum |
| Description pir: [LN:C71375] [AC:C71375] [protein TP0033] [GN:TP0033] [GS:C71375] [syphilis spirochete] [DB:pir2] [AC:AE001188:AE000520] [PN:content [GN:TP0033] [OR:Treponema pall section 4 of 87 of the complete [PID:1001643 percent] [LE:7016] | OR:Trepo >gp:[GI served h idum] [I e genome [RE:762 | erved hyponema pal ::g332228 hypotheti DB:genper :][NT:s :7][DI:c | pothet llidum 38] [L ical i ical i ict-bct simila comple | ical i subsp N:AE00 ntegra 2] [DE r to P ment] | . pallio 1188] 1 membra :Trepona ID:10016 | dum] [SR:, ane] ema pallidum 513 |
| Description pir: [LN:C71375] [AC:C71375] [protein TP0033] [GN:TP0033] [GS:C71375] [syphilis spirochete] [DB:pir2] [AC:AE001188:AE000520] [PN:cont [GN:TP0033] [OR:Treponema pall section 4 of 87 of the complete PID:1001643 percent] [LE:7016] ORF Name | OR:Trepo >gp:[GI served h idum] [I e genome [RE:762 | erved hyponema pal ::g332228 hypotheti DB:genper :][NT:s :7][DI:c | pothet llidum 38] [L ical i ot-bct simila comple NT LN | ical i subsp N:AE00 ntegra 2] [DE r to P ment] | . pallic 1188] 1 membra :Trepone ID:10016 | dum] [SR:, ane] ema pallidum 513 P-Value |
| Description pir: [LN:C71375] [AC:C71375] [protein TP0033] [GN:TP0033] [syphilis spirochete] [DB:pir2] [AC:AE001188:AE000520] [PN:confign:TP0033] [OR:Treponema pallisection 4 of 87 of the complete PID:1001643 percent] [LE:7016] ORF Name AI7503001001_39818_f2_530 | OR:Trepo >gp:[GI served h idum] [I e genome [RE:762 NT ID 3594 [AC:U57 :genpept | erved hyponema pal ::g332228 hypotheti DB:genper ::] [NT:s :7] [DI:c | pothet llidum 38] [L ical i ot-bct simila comple NT LN 240 V:ScdA [DE:St | ical i subsp N:AE00 ntegra 2] [DE r to P ment] AA LN 79] [GN:aphylo | Score 234 scdA coccus a | num] [SR:, ane] ema pallidum 513 P-Value 1.2e-19 |
| Description pir:[LN:C71375] [AC:C71375] [Interpretation of the complete of th | OR:Trepo >gp:[GI served h idum] [I e genome [RE:762 NT ID 3594 [AC:U57 :genpept reus cel | erved hyronema pal ::g332228 hypotheti DB:genper :] [NT:s :7] [DI:c AA ID [7366] [PN :-bct1] [:1s conta | oothet llidum 38] [L ical i ot-bct simila comple NT LN 240 V:ScdA [DE:St aining | ical i subsp N:AE00 ntegra 2] [DE r to P ment] AA LN [GN: aphylo a scd | Score 234 scdA coccus a | num] [SR:, ane] ema pallidum 513 P-Value 1.2e-19 |
| Description pir:[LN:C71375] [AC:C71375] [Incomplete of TP0033] [GN:TP0033] [GN:TP0033] [GN:TP0033] [GN:TP0033] [GN:TP0033] [GN:TP0033] [GN:Treponema pallowed of State of the complete of Tp0033] [Gn:Treponema pallowed of State of the complete of Tp0033] [Gn:Treponema pallowed of State of the complete of Tp0033] [Gn:Treponema pallowed of State of Tp0033] [LE:7016] ORF Name AI7503001001_39818_f2_530 Description gp:[GI:g1575061] [LN:SAU57060] [GR:Staphylococcus aureus] [DB gene, complete cds.] [NT:S. au: [LE:361] [RE:1035] [DI:direct] | OR:Trepo >gp:[GI served h idum] [I e genome [RE:762 NT ID 3594 [AC:U57 :genpept reus cel | erved hyronema pal i:g332228 hypotheti DB:genper e.] [NT:s 7] [DI:c AA ID 7366 | oothet liidum 38] [L ical i ot-bct simila comple NT LN 240 V:ScdA [DE:St aining | ical i subsp N:AE00 ntegra 2] [DE r to P ment] AA LN 79] [GN: aphylo a scd | scdA coccus a disrup | ema pallidum 513 P-Value [1.2e-19 aureus scdA ption have] |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
|---|--|---|---|--|---|---|--|--|
| AI7503001001_4023518_f2_400 | 3596 | 7368 | 909 | 302 | 579 | 3.3e-56 | | |
| Description | | | 1 | , | J | | | |
| pir:[LN:G69800] [AC:G69800] [EGN:yfhF] [CL:cell division in [DB:pir2] >gp:[GI:e1182841:g26] [GN:yfhF] [FN:unknown] [OR:Bac:subtilis complete genome (sect:[NT:similar to cell-division in [DI:complement] >gp:[GI:d102538] [OR:Bacillus subtilis] [SR:Bac:[DE:Bacillus subtilis] [SR:Bac:[LE:14211] [RE:15122] [DI:complement] | nhibitor 33175] [illus su ion 5 of nhibitor 88:g2804 illus su ome sequ | yfhF] [LN:BSUB btilis] [21): f [LE:1 536] [L btilis | [OR:Bac 0005] [DB:go rom 80: 21320] N:D850: DNA] [1 | cillus [AC:Z9: enpept 2821 to [RE:1: 82] [AG DB:gen] | subtili 9108:ALC -bctl] (01011250 22231] C:D85082 pept-bct | [S] 009126] [DE:Bacillus 0.] 2] [PN:YfhF] 3:1] | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503001001_4036093_c3_1384 | 3597 | 7369 | 864 | 287 | 461 | 1.0e-43 | | |
| Description pir: [LN:B69866] [AC:B69866] [PN:transcription regulator LysR family homolog ykuM] [GN:ykuM] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1181918:g2632234] [LN:BS16829KB] [AC:AJ222587] [PN:YkuM protein] [GN:ykuM] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 29kB DNA fragment from ykwC gene to cse15 gene.] [NT:homologous to LysR type transcriptional regulators] [LE:20403] [RE:21284] [DI:direct] >gp:[GI:e1185004:g2633785] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ykuM] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 | | | | | | | | |
| of 21): from 1394791to 1603020 (LysR family)] [LE:90659] [RE: | | | | nscript | cional r | egulator | | |
| ORF Name AI7503001001_4072006_f3_845 Description | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> 201 | AA LN | Score | <u>P-Value</u> | | |

| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | P-Value | | | |
|--|---|---|--|--|---|------------------------------------|--|--|--|
| AI7503001001_4095055_f2_463 | 3599 | 7371 | 894 | 297 | 512 | 4.1e-49 | | | |
| Description | | | | | | | | | |
| pir:[LN:B71256] [AC:B71256] [Fortier of the protein TP0986] [GN:TP0986] [CS:philis spirochete] [DB:pir2] [AC:AE001266:AE000520] [PN:consection 82 of 87 of the complete PID:2314395 percent] [LE:1967] | OR:Trepo >gp:{GI served h ldum] [D ce genom | nema pa :g33233 ypothet B:genpe e.] [NT | llidum 11] [L1 ical in pt-bct2 :simila | subsp N:AE00 ntegra 2] [DE ar to | . pallid 1266] l membra :Trepone | dum] [SR:, nne] ema pallidum | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | | |
| AI7503001001_4096093_c3_1496 | 3600 | 7372 | 354 | 117 | 155 | 2.8e-11 | | | |
| Description | | | | | | | | | |
| pir:[LN:B42573] [AC:B42573] [PN:urf120] [OR:Paracoccus denitrificans] [DB:pir2] | | | | | | | | | |
| ODE Name | NIII TD | 77 TD | NT | AA | | D. Wales | | | |
| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value | | | |
| AI7503001001_4100093_c3_1419 | 3601 | 7373 | 1161 | 386 | 489 | 1.1e-46 | | | |
| Description | | | | | | | | | |
| pir:[LN:E64556] [AC:E64556] [PN:para-aminobenzoate synthetase] [OR:Helicobacter pylori] [DB:pir2] >gp:[GI:g2313385] [LN:AE000547] [AC:AE000547:AE000511] [PN:para-aminobenzoate synthetase (pabB)] [GN:HP0293] [OR:Helicobacter pylori 26695] [DB:genpept-bct2] [DE:Helicobacter pylori 26695 section 25 of 134 of the complete genome.] [NT:similar to GB:K02673 SP:P05041 GB:U07748 GB:U07749] [LE:20379] [RE:22058] [DI:direct] | | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | | |
| AI7503001001_4101643_c2_1314 | 3602 | 7374 | 399 | 132 | 304 | 4.5e-27 | | | |
| Description | | | | | | | | | |
| Description pir:[LN:F70023] [AC:F70023] [PN:hypothetical protein yutD] [GN:yutD] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184310:g2635728] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yutD] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420.] [LE:121344] [RE:121652] [DI:complement] | | | | | | | | | |

TNAA NT ID AA ID ORF Name Score P-Value LN LN AI7503001001 4103393_c2_1215 7375 3603 597 198 436 4.7e-41

Description

sp:[LN:PABA_SALTY] [AC:P06193] [GN:PABA] [OR:SALMONELLA TYPHIMURIUM]
[EC:4.1.3.-] [DE:(EC 4.1.3.-) (ADC SYNTHASE)] [SP:P06193] [DB:swissprot]
>pir:[LN:S09636] [AC:S09636] [PN:pabA protein] [GN:pabA] [CL:glutamine amidotransferase:trpG homology] [OR:Salmonella typhimurium] [DB:pir2]
>gp:[GI:g47816] [LN:STPABA] [AC:X02603] [OR:Salmonella typhimurium]
[DB:genpept-bct1] [DE:Salmonella typhimurium pabA gene for para-aminobenzoate synthaseglutamine amidotransferase.] [NT:pabA gene product (aa 1-187)] [SP:P06193] [LE:1] [RE:564] [DI:direct] >gp:[GI:g154228] [LN:STYPABAA] [AC:M32355] [OR:Salmonella typhimurium] [SR:S.typhimurium DNA, clone pSZD3] [DB:genpept-bct1] [DE:S.typhimurium glutamine amidotransferase subunit ofpara-aminobenzoate synthase (pabA) and pot. cell division proteinfic) genes, complete cds.] [NT:glutamine amidotransferase] [LE:970] [RE:1533] [DI:direct]

NT AΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001 4110888_c3_1380 3604 7376 405 134 93 0.00010

Description

gp:[GI:g3283053] [LN:AF054173] [AC:AF054173] [PN:staphylococcal accessory
regulator A homolog] [GN:sarA] [OR:Staphylococcus epidermidis]
[DB:genpept-bct2] [DE:Staphylococcus epidermidis staphylococcal accessory
regulator Ahomolog (sarA) gene, complete cds.] [LE:887] [RE:1261]
[DI:direct]

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 AI7503001001_4111691_c1_1018
 3605
 7377
 744
 247
 351
 1.5e-31

Description

sp:[LN:YCSJ BACSU] [AC:P42967] [GN:YCSJ] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 63.8 KD PROTEIN IN SIPU-PBPC INTERGENIC REGION] [SP:P42967] [DB:swissprot] >pir:[LN:G69765] [AC:G69765:I39898] [PN:allophanate hydrolase homolog ycsJ:probable urea amidolyase] [GN:ycsJ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1007939:g790943] [LN:BAC39R] [AC:D38161] [PN:urea amidolyase] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168TrpC2) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genome around 39 degrees region encoding 17 ORFs, complete cds.] [LE:10358] [RE:12094] [DI:direct] >gp:[GI:e1182375:g2632709] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:ycsJ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:similar to allophanate hydrolase] [SP:P42967] [LE:56667] [RE:58403] [DI:direct] >gp:[GI:d1009674:g1805477] [LN:D50453] [AC:D50453] [PN:homologues to hypothetical protein HI1731 of H.] [GN:ycsJ] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168 trpC2) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for 25-36 degree region containing theamyE-srfA region, complete cds.] [LE:139065] [RE:140801] [DI:direct]

| ORF Name | NT ID | AA ID | <u>NT</u> LN | AA LN | Score | P-Value |
|-----------------------------|-------|-------|-----------------|----------|-------|----------|
| AI7503001001_4195817_c1_955 | 3606 | 7378 | 1743 | 580 | 2597 | 4.7e-270 |

Description

gp:[GI:e264711:g1262136] [LN:SAPBP4GEN] [AC:X91786] [PN:ATP-binding cassette
transporter A] [GN:abcA] [OR:Staphylococcus aureus] [DB:genpept-bct1]
[DE:S.aureus abcA, pbp4, and tagD genes.] [LE:311] [RE:2038] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|---------------------------|-------|-------|----------|-----------------|-------|----------|
| AI7503001001_42167_c1_959 | 3607 | 7379 | 834 | 277 | 1083 | 1.3e-109 |

Description

gp:[GI:g4928292] [LN:AF132117] [AC:AF132117] [PN:FhuA] [GN:fhuA]
[OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus
ferrichrome uptake operon, complete sequenceand unknown genes.] [NT:ATP
binding protein.] [LE:2780] [RE:3514] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|---|---|--|--------------------------|----------------------------------|--------------------------------------|
| AI7503001001_422162_f3_795 | 3608 | 7380 | 456 | 151 | 439 | 2.2e-41 |
| Description | | | J L — | | | |
| pir:[LN:B69868] [AC:B69868] [A | acillus :BSUB000 ilis] [I 21): fro | subtili 08] [AC: 0B:genpe om 13947 | .s] [DB :Z99111 :pt-bct: /91to 16 | pir2] AL009 L] [DE | 126] [GN :Bacillu .] [NT:s | J:ykvK] us subtilis similar to |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_422800_f3_903 Description | 3609 | 7381 | 447 | 148 | 627 | 2.7e-61 |
| gp:[GI:g3283053] [LN:AF054173] regulator A homolog] [GN:sarA] [DB:genpept-bct2] [DE:Staphylog regulator Ahomolog (sarA) gene [DI:direct] | [OR:Sta | aphyloco epidermi | occus er dis sta | oiderm aphylo | idis] coccal a | accessory |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_425927_f1_232 | 3610 | 7382 | 144 | 47 | | |
| Description NO-HIT | | | | | | |
| ORF Name AI7503001001 429675 f2 558 | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> | Score | P-Value |
| Description | | | الستال | | | |
| NO-HIT | | | / | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_4330390_c3_1517 | 3612 | 7384 | 876 | 291 | 787 | 3.0e-78 |
| Description | | | | | | |
| <pre>pir:[LN:A70016] [AC:A70016] [I [OR:Bacillus subtilis] [DB:pir2 [AC:Z99120:AL009126] [GN:yunF] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [LE:12</pre> | ?] >gp: [FN:un} subtili | [GI:e118 nown] [s compl | 4318:g2 OR:Baci ete ger | 2635730 .llus : | [LN:E subtilis section | SUB0017] |

| ORF Name | NT ID | AA ID | NT | AA | Score | P-Value |
|--|---------------------------------|--------------------------------|----------------------------|------------------------------|----------------------|---------------------|
| A17503001001 4351465 c2 1159 | 3613 | 7385 | <u>LN</u> 1129 | <u>LN</u> 42 | ¬ | |
| Description | |][|][| | | |
| NO-HIT | | | | | | |
| | | | | | - | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_4461693_f2_596 | 3614 | 7386 | 240 | 79 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_4487588_f3_797 | 3615 | 7387 | 609 | 202 | 81 | 0.00094 |
| oxidoreductase, 43.2 kDa subun: [DB:genpept-bct2] [DE:Archaeog: complete genome.] [NT:similar t [LE:4241] [RE:5455] [DI:direct] ORF Name | lobus fu to PID:8 | ılgidus | section | n 131 73 per <u>AA</u> | of 172 c | of the |
| AI7503001001_4501250_c1_1097 | | | LN | <u>LN</u> | | |
| Description | 3616 | 7388 | 1080 | 359 | 871 | 3.7e-87 |
| gp:[GI:d1020364:g1944409] [LN:I protein] [OR:Bacillus stearothe (strain:TRBE14) DNA] [DB:genperglycogen operon genes, complete Alkaligenes eutrophus] [LE:144] | ermophil pt-bct1] e cds.] | us] [SR. [DE:Ba [NT:The] | :Bacil cillus ORF is | lus st stear s simi | earother othermop | cmophilus philus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_4511550_c2_1247 | 3617 | 7389 | 1278 | 425 | 318 | 1.5e-28 |
| Description | | | | | | |
| <pre>sp:[LN:YHAD_ECOLI] [AC:P23524] [DE:HYPOTHETICAL 39.1 KD PROTE] [SP:P23524] [DB:swissprot]</pre> | | | | | | (ORF 3)] |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|---|---------------------------------|------------------------------------|-------------------------------|-------------------------|--------------|---------------|
| A17503001001_4535652_f1_255 Description | 3618 | 7390 | 225 | 74 |] | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_4539143_c1_1066 Description | 3619 | 7391 | 342 | 313 | 970 | 1.2e-97 |
| sp:[LN:HPRK_BACSU] [AC:034483] [DE:HPR(SER) KINASE,] [SP:0344 | | | | SUBT | TILIS] [| EC:2.7.1] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_4578956_f1_101 Description | 3620 | 7392 | 297 | 8 | 79 | 0.021 |
| gp:[GI:e1350598:g3881046] [LN: [OR:Caenorhabditis elegans] [D cosmid Y51A2D, complete sequen [RE:83208:83768:84978] [DI:com | B:genpep .ce.] [LE | t-inv1] :83074:83 | [DE:Cae | norha | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_4687843_c2_1324 Description | 3621 | 7393 3 | 30 | .09 | 242 | 1.7e-20 |
| pir:[LN:C70026] [AC:C70026] [OR:Bacillus subtilis] [DB:pir [AC:Z99120:AL009126] [GN:yuzD] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [LE:1 | 2] >gp:[[FN:unk: subtili | GI:e1184 nown] [OI s complet | 300:g26 R:Bacil ce geno | 35718 lus s me (s | [LN:Bubtilis | SUB0017]] |

NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503001001_4689077_c2_1271 3622 7394 333 110 466 3.1e-44 Description sp:[LN:CLPP BACSU] [AC:P80244:008433] [GN:CLPP] [OR:BACILLUS SUBTILIS] [EC:3.4.21.92] [DE:(ENDOPEPTIDASE CLP) (CASEINOLYTIC PROTEASE) (PROTEASE TI)] [SP:P80244:O08433] [DB:swissprot] >pir:[LN:B69601] [AC:B69601:A47683] [PN:ATP-dependent clp proteinase, chain P:stress protein G7] [GN:clpP] [CL:ATP-dependent Clp proteinase chain P] [OR:Bacillus subtilis] [EC:3.4.21.-] [DB:pir2] >qp:[GI:e1186142:q2635967] [LN:BSUB0018] [AC:Z99121:AL009126] [PN:ATP-dependent Clp protease proteolytic subunit] [GN:clpP] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.4.21.92] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:alternate gene name: yvdN] [SP:P80244] [LE:145744] [RE:146337] [DI:direct] >gp:[GI:e313044:g1945673] [LN:BSZ94043] [AC:Z94043] [PN:hypothetical protein] [GN:yvdN] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic DNA fragment (88 kb).] [NT:similar to CLPP ECOLI ATP-dependent clp protease] [SP:P80244] [LE:35334] [RE:35927] [DI:complement] >gp:[GI:g2668494] [LN:BSU59754] [AC:U59754] [PN:ClpP] [GN:clpP] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis Clp protease proteolytic component (clpP) gene, complete cds.] [NT:proteolytic component of Clp protease] [LE:140] [RE:733] [DI:direct] NT AAORF Name NT ID AA ID Score P-Value LN LN AI7503001001_4689130_f2_614 7395 3623 306 101 81 0.029 Description pir: [LN:A58932] [AC:A58932] [PN:cytochrome C-type biogenesis protein CCMF] [GN:yejR:ccmF] [OR:mitochondrion Cyanidioschyzon merolae] [DB:pir2] >gp:[GI:d1037513:g4115789] [LN:D89861] [AC:D89861] [PN:cytochrome C-type biogenesis protein CCMF] [GN:yejR or ccmF] [OR:Mitochondrion Cyanidioschyzon merolae] [SR:Cyanidioschyzon merolae (strain:10D) mitochondrion DNA] [DB:genpept-pln1] [DE:Cyanidioschyzon merolae mitochondrial DNA, complete sequence.] [LE:16296] [RE:18158] [DI:complement] NTAΑ ORF Name NT ID AA ID Score P-Value LN LNAI7503001001 4695293 c3 1487 3624 7396 171 56 Description

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
|--|----------|--------|----------|-----------------|----------|----------------|--|--|
| A17503001001_4703180_c1_1041 | 3625 | 7397 | 453 | 150 | 541 | 3.5e-52 | | |
| Description | | · | | | | | | |
| gp:[GI:e1393149:g4490608] [LN: [OR:Staphylococcus aureus] [DB ribonucleotide reductase operor | :genpept | -bct1] | [DE:St | aphylo | coccus a | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| AI7503001001_4709575_c2_1309 | 3626 | 7398 | 354 | 117 | | | | |
| <u>Description</u> | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_4726462_f2_572 | 3627 | 7399 | 168 | 55 | | | | |
| <u>Description</u> | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | <u>P-Value</u> | | |
| AI7503001001_4726527_f3_774 | 3628 | 7400 | 528 | 175 | 749 | 3.2e-74 | | |
| Description | | | | | | | | |
| pir:[LN:D69868] [AC:D69868] [PN:conserved hypothetical protein ykvM] [GN:ykvM] [CL:hypothetical protein ykvM] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184965:g2633746] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ykvM] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:similar to hypothetical proteins] [LE:45967] [RE:46464] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| AI7503001001_4773392_f2_359 | 3629 | 7401 | 288 | 95 | | | | |
| Description | | | | | | | | |
| | | | | | | | | |

| ORF Name | NT ID | AA ID | $\underline{\mathtt{NT}}$ | <u>AA</u> | Score | P-Value |
|--|---|---|---|---|---|--|
| | | | <u>LN</u> | <u>LN</u> | | |
| A17503001001_4775287_c3_1420 | 3630 | 7402 | 1008 | 335 | 238 | 1.3e-34 |
| Description | | | | | | |
| sp:[LN:YBGK_ECOLI] [AC:P75745] [DE:HYPOTHETICAL 34.4 KD PROTE: [DB:swissprot] >pir:[LN:G64806] [CL:hypothetical protein HI1730] >gp:[GI:d1036362:g4062308] [LN:protein HI1730] [OR:Escherichia:clone:Kohara clone #175] [DB:gettlene:Kohara clone #175] [DB:ge | IN IN PH [AC:G6 [OR:E D90710] coli] enpept-b col75#7; >gp:[G ive carb cherichi | RB-NEI 4806] scheric [AC:D9 [SR:Esclet1] [D] simila I:g1786 oxylase a coli] on 64 of | INTERGH [PN:ybo hia col 0710:AF herichi E:Esche r to Pl 930] [I [GN:y [DB:ge f 400 co | ENIC RESERVED TO THE PROPERTY OF THE PROPERTY | EGION] [cein] [Gein] [Gein] [PN:Hi(strain a coli gession Noo174] [FN:puta-bct2] complet | N:ybgK] ypothetical :K12) DNA, enomic DNA. umber] tive egenome.] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_47763_f3_814 | 3631 | 7403 | 186 | 61 |] | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_4804643_f1_90 | 3632 | 7404 | 246 | 81 | 75 | 0.0026 |
| Description | | | | | | |
| <pre>gp:[GI:e1332553:g3764009] [LN:F [OR:Plasmodium falciparum] [SR: [DB:genpept-inv1] [DE:Plasmodium] [NT:predicted using hexExon; MF [DI:direct]</pre> | malaria ım falci | parasit | ce P. f AL3P4, | alcipa comple | arum] ete seque | ence.] |

NTORF Name AA ID NT ID Score P-Value LN LN 7405 AI7503001001_4806575_c2_1171 3633 651 216 503 3.7e-48 Description

pir:[LN:H69854] [AC:H69854] [PN:hypothetical protein ykaA] [GN:ykaA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1181486:g2632006] [LN:BSAJ2571] [AC:AJ002571] [PN:YkaA] [GN:ykaA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 56 kb DNA fragment between xlyA and ykoR.] [LE:2613] [RE:3230] [DI:complement] >gp:[GI:e1183305:g2633639] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:ykaA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7

 ORF Name
 NT ID
 AA ID
 NT LN
 AA LN
 Score
 P-Value

 AI7503001001_4876077_f3_796
 3634
 7406
 714
 237
 778
 2.7e-77

of 21): from 1194391to 1411140.] [LE:155562] [RE:156179] [DI:complement]

Description

pir:[LN:C69868] [AC:C69868] [PN:coenzyme PQQ synthesis homolog ykvL]
[GN:ykvL] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184964:g2633745]
[LN:BSUB0008] [AC:Z99111:AL009126] [GN:ykvL] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:similar to coenzyme PQQ synthesis]
[LE:45218] [RE:45949] [DI:direct]

NT AΑ ORF Name NT ID AA ID Score P-Value LN LN A17503001001_4885876_c1_1071 918 305 3635 7407 829 1.1e-82

Description

sp:[LN:YVCJ_BACSU] [AC:006973] [GN:YVCJ] [OR:BACILLUS SUBTILIS]
[DE:HYPOTHETICAL 33.9 KD PROTEIN IN CRH-TRXB INTERGENIC REGION] [SP:006973]
[DB:swissprot] >pir:[LN:H70031] [AC:H70031] [PN:conserved hypothetical
protein yvcJ] [GN:yvcJ] [CL:Bacillus subtilis conserved hypothetical
protein yvcJ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1186165:g2635990]
[LN:BSUB0018] [AC:Z99121:AL009126] [GN:yvcJ] [FN:unknown] [OR:Bacillus
subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section
18 of 21): from 3399551to 3609060.] [NT:similar to hypothetical proteins]
[SP:006973] [LE:171011] [RE:171898] [DI:complement]
>gp:[GI:e313026:g1945650] [LN:BSZ94043] [AC:Z94043] [PN:hypothetical
protein] [GN:yvcJ] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis
genomic DNA fragment (88 kb).] [NT:similar to hypothetical MTCY21B4]
[SP:006973] [LE:9773] [RE:10660] [DI:direct]

[RE:420] [DI:complement]

| ORF Name | NT ID | AA ID | , <u>NT</u> | AA | Score | P-Value |
|---|---------------------------------|----------------------------------|-------------------------------|-----------------------------|--------------------------------|---------------------|
| AI7503001001 4892878 c3 1357 | 13636 | 17408 | <u>LN</u> (1077 | <u>LN</u> 358 | 1455 | 4.5e-43 |
| Description | | | | | التقيا | |
| gp:[GI:g1913906] [LN:SAU91741] acid biosynthesis] [OR:Staphylo [DE:Staphylococcus aureus teich and TagX and TagD genes, comple | ococcus hoic aci | aureus] d biosy | DB:ge nthesis | enpept s TagB | -bct1] gene, p | partialcds |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_4898376_f2_574 | 3637 | 7409 | 207 | 68 | 52 | 0.024 |
| Description | | | | | | |
| <pre>gp:[GI:g4588017] [LN:AF087657] oxidoreductase chain 6] [GN:ND6 [SR:Agrocybe aegerita] [DB:geng NADH-ubiquinone oxidoreductase encoding mitochondrial protein, [LE:34] [RE:453] [DI:direct]</pre> | 6] [OR:M pept-pln chain 6 | Mitochon n2] [DE: 5 (ND6)g | drion A Agrocyl ene, mi | Agrocy be aego itocho | be aeger erita ndrial g | , jene |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_5084652_f3_849 | 3638 | 7410 | 507 | 168 | 536 | 1.2e-51 |
| Description | | | | | | |
| <pre>gp:[GI:g1575061] [LN:SAU57060] [OR:Staphylococcus aureus] [DB:gene, complete cds.] [NT:S. aur [LE:361] [RE:1035] [DI:direct]</pre> | :genpept | -bct1] | [DE:Sta | aphylo | coccus a | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_5117793_c3_1531 | 3639 | 7411 | 273 | 91 | 197 | 9.9e-16 |
| Description sp:[LN:YRN3_VIBPA] [AC:P46231] PROTEIN IN RNT 5'REGION (ORF3) >gp:[GI:g497126] [LN:VPU06949] parahaemolyticus] [DB:genpept-k (rnt) gene and flagellar motore | FRAGME) AC:U06 oct1] [D | NT)] [S: 949] [P: E:Vibri | P:P4623 N:ORF3] o parah | 31] [DI OR:V naemoly | B:swissp Vibrio Yticus B | rot] B22 RNase T |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
|--|----------|---------------------|----------|--------------------|----------------------|-----------------------|--|--|
| AI7503001001_5131265_c2_1243 | 3640 | 7412 | | 91 | 80 | 0.0071 | | |
| Description | <u> </u> | JI | l L | J L | J [] | · | | |
| <pre>gp:[GI:g3582235] [LN:AE001272] region ORF00014] [GN:ORF00014] [DE:Lactococcus lactis DPC3147 [NT:hypothetical protein; ident [DI:direct]</pre> | [OR:Lac | tococcu 1 pMRC01 | s lact: | is] [D] lete p: | B:genpep lasmidse | ot-bct2] equence.] | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_5182962_f1_196 Description | 3641 | 7413 | 1068 | 355 | 295 | 1.3e-30 | | |
| sp:[LN:RESE_BACSU] [AC:P35164] [GN:RESE] [OR:BACILLUS SUBTILIS] [EC:2.7.3] [DE:SENSOR PROTEIN RESE,] [SP:P35164] [DB:swissprot] >pir:[LN:H69691] [AC:H69691:S45560] [PN:two-component sensor histidine kinase resE] [GN:resE] [CL:sensor histidine kinase homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g410142] [LN:BACDIA] [AC:L09228] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain 168, sub_species Marburg) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis spoVA to serA region.] [NT:ORFX18] [LE:22425] [RE:24194] [DI:direct] >gp:[GI:e1185580:g2634746] [LN:BSUB0013] [AC:Z99116:AL009126] [PN:two-component sensor histidine kinase] [GN:resE] [FN:involved in global regulation of aerobic and] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:alternate gene name: ypxE] [SP:P35164] [LE:19389] [RE:21158] [DI:complement] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503001001_5195393_c3_1352 | 3642 | 7414 | 654 | 217 | 1088 | 3.8e-110 | | |
| Description | | | | | | | | |
| <pre>gp:[GI:e255543:g1617434] [LN:SE repressor] [GN:sirR] [OR:Staphy [DE:S.epidermidis sirR gene.] [</pre> | /lococcu | s epide: | rmidis] | [DB:g | genpept- | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
|---|---|---|--|---|---------------------------------|----------|--|--|
| A17503001001_5276677_c1_1026 | 3643 | 7415 | 1566 | 521 | 703 | 2.4e-69 | | |
| Description | | | | <u> </u> | | | | |
| pir:[LN:B64622] [AC:B64622] [PN:osmoprotection protein] [OR:Helicobacter pylori] [DB:pir2] >gp:[GI:g2313949] [LN:AE000593] [AC:AE000593:AE000511] [PN:osmoprotection protein (proWX)] [GN:HP0818] [OR:Helicobacter pylori 26695] [DB:genpept-bct2] [DE:Helicobacter pylori 26695 section 71 of 134 of the complete genome.] [NT:similar to PID:1109685 SP:Q45461 GB:AL009126] [LE:4862] [RE:6523] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_5283592_f1_215 | 3644 | 7416 | 288 | 95 | 73 | 0.014 | | |
| Description | | | | | _ | | | |
| <pre>gp:[GI:g1054677] [LN:CCCOX3] [AC:X92734] [PN:cytochrome oxidase] [GN:coxIII] [OR:Mitochondrion Chara corallina] [SR:Chara corallina] [DB:genpept-pln1] [DE:C.corallina mitochondrial cox3 gene.] [NT:subunit III] [LE:<1] [RE:>381] [DI:direct]</pre> | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_5292175_c3_1484 | 3645 | 7417 | 1323 | 440 | 2112 | 1.2e-218 | | |
| Description | | | | | | | | |
| gp:[GI:g3152725] [LN:AF065394] [AC:AF065394] [PN:enolase] [GN:eno] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus enolase (eno) gene, complete cds.] [NT:ENO; laminin binding protein] [LE:103] [RE:1407] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT | AA | Score | P-Value | | |
| | | | LN | LN | | | | |
| A17503001001_53552_f2_450 | 3646 | 7418 | 660 | 219 | 179 | 8.0e-14 | | |
| Description sp:[LN:GRPB_BACFI] [AC:Q45133] [DE:GLUTAMATE-RICH PROTEIN GRPF >gp:[GI:g1209681] [LN:BFU39410] [GN:grpB] [OR:Bacillus firmus] [DB:genpept-bct1] [DE:Bacillus (grpA), OrfC, and glutamate-rich [LE:2695] [RE:3219] [DI:direct] | 3] [SP:Q [AC:U3 SR:Bac firmus n protei | 45133] 9410] [illus f OrfA, O | [DB:swi PN:glut irmus s rfB, gl | issprot amate- strain= lutamat | c] rich pr OF4] e-rich | protein | | |

 ORF Name
 NT ID
 AA ID
 NT ID
 AA ID
 NT ID
 AA ID
 NT ID
 AA ID
 LN ID
 LN ID
 Score
 P-Value

 AI7503001001_5355325_c3_1454
 | 3647 | 7419 | 888 | 295 | | 526 | | 1.4e-50 | |
 | 1.4e-50 | |
 | 1.4e-50 | |
 | 1.4e-50 | |
 | 1.4e-50 | |
 | 1.4e-50 | |
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 | 1.4e-50 | |
 | 1.4e

Description

sp:[LN:DEGV BACSU] [AC:P32436] [GN:DEGV] [OR:BACILLUS SUBTILIS] [DE:DEGV PROTEIN] [SP:P32436] [DB:swissprot] >pir:[LN:D30191] [AC:I40386:D30191:A70042:S28596] [PN:conserved hypothetical protein yviA] [GN:yviA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g39848] [LN:BSCOMFG] [AC:Z18629] [PN:U3] [GN:degUorf3] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis comF gene.] [SP:P32436] [LE:1] [RE:846] [DI:direct] >gp:[GI:e1184454:g2636074] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:yviA] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: degV, sacU; similar to] [LE:45634] [RE:46479] [DI:complement] >gp:[GI:g1762331] [LN:BSU56901] [AC:U56901] [GN:degV] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis putative transcriptional regulator (yvhJ), Ycr59c/YigZ homolog (yvhK), histidine kinase (degS), transcriptional regulator of degradation enzyme (degU), (degV), (comFA), (comFB), (comFC), flagellar protein (yviB), negative regulator of flagellin(flgM), flagellar protein (yviC), flagellar-hook associated protein1 (flgK), flagellar-hook associated protein 3 (flgL), (yviE), transmembrane protein (yviF), (csrA), flagellin (hag), flagellarprotein (yviH), flagellar hook-associated protein 2 (fliD), flagellar protein (fliS), flagellar protein (fliT), sigma-54modulator homolog (yviI), and (secA) genes, complete cds.] [LE:4114] [RE:4959] [DI:direct] >gp:[GI:e1184454:g2636074] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:yviA] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: degV, sacU; similar to] [SP:P32436] [LE:45634] [RE:46479] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|------------------------------|-------|-------|----------|----------|-------|---------|
| AI7503001001_5367843_c3_1488 | 3648 | 7420 | 759 | 252 | 791 | 1.1e-78 |
| Dan and a bid and | | | | | | |

Description

sp:[LN:EST_BACST] [AC:Q06174] [GN:EST] [OR:BACILLUS STEAROTHERMOPHILUS]
[EC:3.1.1.1] [DE:CARBOXYLESTERASE PRECURSOR,] [SP:Q06174] [DB:swissprot]
>pir:[LN:JC1374] [AC:JC1374] [PN:carboxylesterase,] [GN:Est] [OR:Bacillus stearothermophilus] [EC:3.1.1.1] [DB:pir2] >gp:[GI:d1002674:g216314]
[LN:BACPBH7] [AC:D12681] [PN:esterase] [OR:Bacillus stearothermophilus]
[SR:Bacillus stearothermophilus DNA, clone pBH7] [DB:genpept-bct1]
[DE:Bacillus stearothermophilus esterase gene.] [LE:181] [RE:924]
[DI:direct]

| ORF Name | NT ID | AA ID | NT | AA | Score | P-Value | | |
|--|---------------------------------|--|----------------------------|-----------------------------|----------------------------------|-----------------|--|--|
| A17503001001_57800_c3_1494 | 3649 | 7421 | <u>LN</u> 147 | <u>LN</u> 48 |] | | | |
| Description | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| AI7503001001_5869433_f2_542 | 3650 | 7422 | 129 | 42 |] | | | |
| Description | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_5869702_f2_595 | 3651 | 7423 | 309 | 102 | 87 | 0.015 | | |
| Description | | | | | | | | |
| sp:[LN:YMW6_YEAST] [AC:Q04264:Q04780] [GN:YMR076C:YM9582.01C:YM9916.15C] [OR:SACCHAROMYCES CEREVISIAE] [SR:,BAKER'S YEAST] [DE:HYPOTHETICAL 147.0 KD PROTEIN IN ABF2-CHL12 INTERGENIC REGION] [SP:Q04264:Q04780] [DB:swissprot] >pir:[LN:S54451] [AC:S54451:S52836] [PN:hypothetical protein YMR076c:hypothetical protein YM9582.01c:hypothetical protein YM9916.15c] [GN:PDS5] [OR:Saccharomyces cerevisiae] [DB:pir2] [MP:13R] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503001001_5938762_c1_987 | 3652 | 7424 | 1704 | 567 | 566 | 7.8e-55 | | |
| Description | | | | | | | | |
| sp:[LN:CYDD_HAEIN] [AC:P45082] [DE:TRANSPORT ATP-BINDING PROTE >pir:[LN:F64186] [AC:F64186]] [CL:unassigned ATP-binding cathomology] [OR:Haemophilus influ | EIN CYDD [PN:ABC- assette | [SP:Petype transported transported to protein the second term of the s | 45082] anspor s: ATP | DB:sv t prote bindir- | wissprot ein cydD ng casse |]] [GN:cydD | | |

[LN:U32795] [AC:U32795:L42023] [PN:ATP-binding protein protein (cydD)] [GN:HI1157] [OR:Haemophilus influenzae Rd] [DB:genpept-bct2] [DE:Haemophilus influenzae Rd section 110 of 163 of the completegenome.] [NT:similar to GB:L21749 SP:P29018 GB:L25859 PID:146416] [LE:6353] [RE:8113] [DI:complement]

| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
|---|--|---|---|--------------------------------------|--|-----------------------------------|
| AI7503001001_6023915_c2_1313 | 3653 | 7425 | 969 | 322 | 1144 | 4.4e-116 |
| Description | | | | | | |
| <pre>pir:[LN:D70023] [AC:D70023] [[CL:lipoic acid synthase] [OR: >gp:[GI:e1184312:g2635730] [LN [FN:unknown] [OR:Bacillus subt complete genome (section 17 of lipoic acid synthetase] [LE:12</pre> | Bacillus [:BSUB001 ilis] [D 21]: fr | s subtil: .7] [AC:2 B:genper :om 31970 | is] [E0 Z99120 pt-bct: 001to : | C:2.8. :AL009 L] [DE 341442 | 1] [DE 126] [GN :Bacillu 0.] [NT: | B:pir2] J:yutB] Is subtilis |
| ORF Name A17503001001_6037756_f3_682 | NT ID | <u>AA ID</u> | NT LN 150 | <u>AA</u> <u>LN</u> 49 | Score | P-Value |
| Description | | l | | <u> </u> | _] | |
| NO-HIT | | | | , | | |
| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> <u>LN</u> | Score | P-Value |
| AI7503001001_6050010_f1_177 | 3655 | 7427 | 132 | 43 | | |
| <u>Description</u> | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_6051500_c1_972 | 3656 | 7428 | 189 | 62 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| | | | | | | |

| ORF Name [AI7503001001 6051537 c1 1063 | 3657 | 7429 | <u>LN</u> 276 | <u>LN</u> 1 191 | <u>Score</u> | 1.0e-06 |
|--|------|------|------------------|--------------------|--------------|---------|
| M17503001001_6051537_C1_1063 | 3657 | 1429 | 276 | 91 | | 1.0e-06 |

Description

sp:[LN:CSBA BACSU] [AC:P37953] [GN:CSBA] [OR:BACILLUS SUBTILIS] [DE:CSBA PROTEIN] [SP:P37953] [DB:swissprot] >gp:[GI:g142780] [LN:BACCSBA] [AC:M80473] [GN:csbA] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain 168, sub species Marburg) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis csbA and uvr/dinA genes, complete cds.] [NT:putative membrane protein; putative] [LE:380] [RE:610] [DI:direct] >gp:[GI:g142842] [LN:BACDINA76] [AC:M64048] [PN:DNase inhibitor] [GN:dinA] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNase inhibitor (dinA76) gene, complete cds andpromoter region.] [LE:64] [RE:294] [DI:direct] >gp:[GI:e1184424:g2636044] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:putative membrane protein] [GN:csbA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P37953] [LE:17086] [RE:17316] [DI:complement] >gp:[GI:g2618840] [LN:AF017113] [AC:AF017113] [PN:CsbA] [GN:csbA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis 300-304 degree genomic sequence.] [LE:12977] [RE:13207] [DI:direct] >gp:[GI:e1184424:g2636044] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:putative membrane protein] [GN:csbA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P37953] [LE:17086] [RE:17316] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|-------|-------|----------|----------|-------|---------|
| A17503001001_6053176_f1_195 | 3658 | 7430 | 519 | 172 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_6056628_c3_1339 Description | 3659 | 7431 | 150 | 49 |] | |
| | | | | | | |

| ORF Name | NT ID | AA ID | LN | LN LN | Score | P-Value |
|--|-------------|---------|-------------------|----------|---------|----------------|
| A17503001001_6057338_c3_1518 | 3660 | 7432 | 840 | 279 | 622 | 9.1e-61 |
| Description | | | | | | |
| <pre>pir:[LN:H70015] [AC:H70015] [[OR:Bacillus subtilis] [DB:pir</pre> | | | - | - | | |
| [AC:Z99120:AL009126] [GN:yunE] | | | _ | | | |
| [DB:genpept-bct1] [DE:Bacillus | | _ | _ | | | 17 of 21): |
| from 3197001to 3414420.] [LE:1 | 2/136] | RE:12/9 | נען ני | .:comp. | rement, | |
| ORF Name | NT ID | AA ID | NT | AA LN | Score | P-Value |
| A17503001001_6094177_c2_1235 | 3661 | 7433 | <u>LN</u> [123 | 40 | 7 | - |
| Description | | | | | _ | |
| NO-HIT | | | | - | | |
| ORF Name | NT ID | AA ID | <u>NT</u> | AA | Score | P-Value |
| | | | LN | LN | 7 | <u>r varac</u> |
| AI7503001001_6101581_f1_74 Description | 3662 | 7434 | 180 | 59 | _ | |
| | | | | | | |
| NO-HIT | | | | | | <u></u> |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_6257763_c3_1479 | 3663 | 7435 | 660 | 219 | 189 | 1.7e-13 |
| Description | | | | 1 | | |

gp:[GI:g2246532] [LN:U93872] [AC:U93872] [OR:Kaposi's sarcoma-associated herpesvirus] [SR:Kaposi's sarcoma-associated herpesvirus - Human herpesvirus 8] [DB:genpept-vrl] [DE:Kaposi's sarcoma-associated herpesvirus glycoprotein M, DNAreplication protein, glycoprotein, DNA replication protein, FLICEinhibitory protein and v-cyclin genes, complete cds, and tegumentprotein gene, partial cds.] [NT:ORF 73, contains large complex repeat CR 73] [LE:124324] [RE:127593] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|--|--|---|---|---|---------------------|
| A17503001001_626592_c1_935 | 3664 | 7436 | 852 | 283 | 233 | 1.5e-19 |
| Description | | | | | | |
| pir:[LN:A69463] [AC:A69463] [Racid hydrolase (pcbD) homolog] >gp:[GI:g2648849] [LN:AE000986] [PN:2-hydroxy-6-oxo-6-phenylhes [OR:Archaeoglobus fulgidus] [DE section 121 of 172 of the compl percent identity: 29.41;] [LE:1 | OR:Arc [AC:AF Ka-2,4-c B:genper Lete ger | chaeoglo 2000986: dienoic ot-bct2] nome.] [1 | bus fu AE0007 acid] [DE:A: NT:sim | lgidus 82] [GN:AF rchaeo ilar t |] [DB:pi 1706] globus f o GP:139 | r2] Tulgidus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_6423376_f3_670 | 3665 | 7437 | 261 | 86 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name AI7503001001_6437525_f3_862 | NT ID | <u>AA ID</u> | NT LN 294 | <u>AA</u> <u>LN</u> 97 | Score | P-Value |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| | | | NTT | 7.7. | • • | **** |
| ORF Name | NT ID | AA ID | $\frac{\text{NT}}{\text{LN}}$ | <u>AA</u> LN | Score | <u>P-Value</u> |
| A17503001001_6440640_c3_1379 | 3667 | 7439 | 1278 | 425 | 139 | 6.0e-06 |
| Description | | | | | | |
| pir:[LN:D71621] [AC:D71621] [ED:D71621] [ED:Plasmodium falciparum] [ED:Plasmodium falciparum] [ED:Plasmodium falciparum] [SR:DB:genpept-inv2] [DE:Plasmodium falciparum] [SR:[DB:genpept-inv2] [DE:Plasmodium falciparum] [NT:plasmodium falc | DB:pir2] othetica malaria um falci | >gp:[G: il prote: parasi: parum cl | I:g384! in] [GI te P. : | 5117] N:PFB0 falcip ome 2, | [LN:AE00 185w] arum] section | 1378] 1 15 of 73 |
| ORF Name | NT ID | AA ID | $\underline{\text{NT}}$ | <u>AA</u> | Score | P-Value |
| A17503001001 6444037 c2 1180 | | | LN | LN | <u> </u> | z varue |
| Description | 3668 | 7440 | 144 | 47 | | |
| NO-HIT | | | | | | |
| 110 1111 | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value | | |
|---|---|---|---|---|---|--|--|--|
| AI7503001001_6642827_c3_1342 | 3669 | 7441 | 2403 | 800 | 1527 | 1.1e-156 | | |
| Description | | · · · · · · · · · · · · · · · · · · · | l <u>— — — — — — — — — — — — — — — — — — —</u> | | | | | |
| pir:[LN:A70010] [AC:A70010] [F [OR:Bacillus subtilis] [DB:pir2 [AC:Z99120:AL009126] [GN:yufT] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:si [RE:51082] [DI:direct] >gp:[GI: [PN:unknown] [GN:yufT] [OR:Baci [DE:B.subtilis genomic DNA frag antiporter/phaA homologue/NADH] | :] >gp:[[FN:unk subtili milar t e311455 llus su | [GI:e118 known] [.s compl to NADH 5:g19348 hbtilis] | 4238:g2 OR:Baci ete ger dehydro 17] [LN [DB:ge to yuf | :635656 .llus : nome (: ogenase J:BSZ9: enpept- EV.] [1 | [LN:Esubtilis section [LE:4 3937] [A -bctl] | SSUB0017] i] 17 of 21): 8758] iC:Z93937] | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| A17503001001_6650312_f2_600 | 3670 | 7442 | 141 | 46 |] | | | |
| Description | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503001001_6675016_c1_1077 | 3671 | 7443 | 132 | 43 |] | | | |
| Description | | | | | | | | |
| NO-HIT | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| AI7503001001_6680312_f2_518 | 3672 | 7444 | 714 | 237 | 211 | 4.7e-17 | | |
| Description | | | | | | | | |
| <pre>gp:[GI:g3329623] [LN:CELF36H12] [AC:AF078790] [GN:F36H12.3] [OR:Caenorhabditis elegans] [DB:genpept-inv2] [DE:Caenorhabditis elegans cosmid F36H12.] [NT:coded for by C. elegans cDNA CEMSF30F] [LE:21606:21872:22500:22677] [RE:21701:22454:22629:22875] [DI:directJoin]</pre> | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503001001_6681577_c1_1110 | 3673 | 7445 | 432 | 143 | 277 | 3.3e-24 | | |
| Description | | | | | | | | |
| pir:[LN:G70020] [AC:G70020] [PN:hypothetical protein yusF] [GN:yusF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184356:g2635774] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yusF] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420] [LE:167450] [RE:167890] [DI:complement] | | | | | | | | |

| ORF Name | NT ID | AA ID | $\frac{\text{NT}}{\text{LN}}$ | AA LN | Score | P-Value | |
|---|---------------------|--------------------|-------------------------------|-------------------------------|--------------------|-----------------|--|
| A17503001001_6695968_c2_1320 | 3674 | 7446 | 1470 | 489 | 2005 | 2.5e-207 | |
| Description gp:[GI:g4530241] [LN:AF101234] protein ligase DltA] [GN:dltA] [DE:Staphylococcus aureus dltAB gene.] [LE:1590] [RE:3047] [DI: | OR:Sta CD oper | phyloco | ccus a | ureus] | [DB:gen | pept-bct2] | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
| AI7503001001_6822175_c2_1222 Description | 3675 | 7447 | 1914 | 637 | 1229 | 4.3e-125 | |
| pir:[LN:F69901] [AC:F69901] [PN:DNA helicase recQ:ATP-dependent DNA helicase homology yocI] [GN:yocI] [CL:recQ protein:DEAD/H box helicase homology:recQ helicase homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g2619051] [LN:AF027868] [AC:AF027868] [PN:RecQ homolog] [GN:yocI] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosome region between terC and odhAB.] [NT:similar to E.coli RecQ protein (607 aa)] [LE:74004] [RE:75779] [DI:complement] >gp:[GI:e1185394:g2634315] [LN:BSUB0011] [AC:Z99114:AL009126] [GN:yocI] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [NT:similar to ATP-dependent DNA helicase] [LE:93095] [RE:94870] [DI:complement] | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
| AI7503001001_6823453_c3_1346 | 3676 | 7448 | 483 | 160 | 270 | 1.8e-23 | |
| Description gp:[GI:d1036085:g4001729] [LN:A [OR:Staphylococcus aureus] [SR: [DB:genpept-bct1] [DE:Staphyloc MnhC, MnhD, MnhE,MnhF and MnhG, [DI:direct] | Staphyl coccus a | ococcus ureus g | aureus enes fo] [LE: | s (stra or OrfA 5474] [| in:209P , MnhA, |) DNA] MnhB, | |
| ORF Name A17503001001_6829638_c3_1402 Description NO-HIT | NT ID | AA ID 7449 | <u>NT</u> <u>LN</u> | AA LN 61 | Score | <u>P-Value</u> | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|--|---|--|---|--|--|
| AI7503001001_6833313_c3_1371 | 3678 | 7450 | 1920 | 639 | 550 | 3.9e-53 |
| Description | | · | L | · | J [] | |
| pir:[LN:B70001] [AC:B70001] [1 [GN:ytsD] [OR:Bacillus subtil: [LN:BSUB0016] [AC:Z99119:AL0093 subtilis] [DB:genpept-bct1] [DI 16 of 21): from 2997771to 32134 (permease)] [LE:110708] [RE:112 [LN:AF008220] [AC:AF008220] [PI [DB:genpept-bct2] [DE:Bacillus [NT:similarity to NADH dehydrog | is] [DB: 126] [GN E:Bacill 410.] [N 2648] [D N:YtsD] subtili | pir2] > (sytsD] us subt T:simil() I:compl() [GN:yts] s rrnB-(| gp:[GI [FN:un] ilis co ar to A ement] D] [OR dnaB ge | ell859 known] omplete ABC tra >gp:[(:Bacill enomic | 910:g263 [OR:Bac e genome ansporte GI:g2293 lus subt region. | s5521] cillus c (section er s178] cilis] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_6836088_c1_1118 | 3679 | 7451 | 1320 | 439 | 1260 | 2.2e-128 |
| Description | | | | | | |
| [GN:yurX] [OR:Bacillus subtile [LN:BSUB0017] [AC:Z99120:AL0099 subtilis] [DB:genpept-bct1] [DB 17 of 21): from 3197001to 34144 [LE:160723] [RE:162036] [DI:con | 126] [GN E:Bacill 420.] [N | :yurX] us subt T:simila | [FN:unlilis co | known] omplete | OR:Bac e genome | cillus e (section |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503001001_6929512_c1_1119 | 3680 | 7452 | 1416 | 471 | 2098 | 3.6e-217 |
| Description | | | | <u> </u> | J [| |
| pir:[LN:D70019] [AC:D70019] [IGN:yurU] [CL:Methanobacterium Ycf24] [OR:Bacillus subtilis] [LN:BSUB0017] [AC:Z99120:AL0091 subtilis] [DB:genpept-bct1] [DB:T of 21): from 3197001to 34144 [LE:157652] [RE:159049] [DI:com | n thermo [DB:pir2 126] [GN E:Bacill 120.] [N | autotrop] >gp:[0 :yurU] us subt: T:simila | phicum GI:e118 (FN:un) ilis co | ABC to 34345:g cnown] omplete | cansport 32635763 [OR:Bac genome | er chain] :illus : (section |
| ORF Name AI7503001001_7042878_f1_40 Description | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> 177 | <u>AA</u> <u>LN</u> 58 | <u>Score</u> | P-Value |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|--|---|---|---|--|---|
| A17503001001_7239188_f2_519 | 3682 | 7454 | 675 | 224 | 491 | 6.9e-47 |
| Description | | | | | | |
| pir:[LN:D70033] [AC:D70033] [3] [GN:yvdD] [CL:yeast conserved subtilis] [DB:pir2] >gp:[GI:el] [AC:Z99121:AL009126] [GN:yvdD] [DB:genpept-bct1] [DE:Bacillus from 3399551to 3609060.] [NT:si [RE:159025] [DI:complement] >gp [AC:Z94043] [PN:hypothetical pictory [DB:genpept-bct1] [DE:B.subtilito YJF5_YEAST hypothetical 26.5] | hypothe 186152:g [FN:unk subtili imilar t p:[GI:e3 rotein] is genom | etical p g2635977 known] [is compl to hypot 313036:g [GN:yvo | protein [OR:Bac: ete ger chetical [1945663] [OR:fragmen | YJL05 BSUB00 illus nome (l prot B] [LN Bacil | 5w] [OR: 18] subtilis section eins] [I :BSZ9404 lus subt kb).] | Bacillus 18 of 21): E:158450] 3] cilis] [NT:similar |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_7800_c1_1093 | 3683 | 7455 | 198 | 65 | 54 | 0.033 |
| Description gp:[GI:g3789915] [LN:AF083442] G2A] [OR:Mus musculus] [SR:hous protein-coupled receptor G2A mI [DI:direct] | se mouse | e] [DB:g | enpept - | rod] | [DE:Mus | musculus G |
| ORF Name AT7503001001_781415_c3_1393 Description | NT ID | <u>AA ID</u> 7456 | NT LN 165 | <u>AA</u> <u>LN</u> 54 | Score | P-Value |
| NO-HIT | | | | | | |
| ORF Name AI7503001001_783375_c2_1236 | NT ID | AA ID | NT LN | <u>AA</u> <u>LN</u> 59 | Score | P-Value |
| Description | | | | | | |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|---|--|--|--|---|---|---|
| A17503001001_788950_c3_1347 | 3686 | 7458 | 531 | 176 | 219 | 4.6e-18 |
| Description | | | | | , | - |
| pir:[LN:E70008] [AC:E70008] [1 [GN:yufB] [OR:Bacillus subtil: [LN:BSUB0017] [AC:Z99120:AL0093 subtilis] [DB:genpept-bct1] [DI 17 of 21): from 3197001to 34144 [LE:54068] [RE:54442] [DI:direction [AC:Z93932] [PN:unknown] [GN:yufube:B.subtilis genomic DNA frag [DI:complement] | is] [DB: 126] [GN E:Bacill 420.] [N ct] >gp: 1fB] [OR | pir2] > :yufB] us subt T:simil [GI:e31 :Bacill | gp:[GI: [FN:unk ilis co ar to h 1512:g1 us subt | e11842 nown] mplete ypothe 934774 | 243:g263 [OR:Bac genome etical p [LN:B [DB:gen | 5661] illus (section roteins] SZ93932] pept-bct1] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_821963_c2_1165 | 3687 | 7459 | 546 | 181 | 103 | 0.0018 |
| Description | | | | - | | - |
| <pre>gp:[GI:g4731376] [LN:AF135127] reductase homolog Bet v 5] [GN:birch] [DB:genpept-pln2] [DE:Bev 5 (BETV5) mRNA,partial cds.] [RE:>900] [DI:direct]</pre> | :BETV5] etula pe | [OR:Bet ndula i | ula pen soflavo | dula] ne red | [SR:Eur luctase | opean white homolog Bet |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_822777_c3_1397 | 3688 | 7460 | 159 | 52 |] | |
| Description | | | | | | |
| NO-HIT | | | | | | <u>'</u> |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_870300_c1_958 | 3689 | 7461 | 153 | 50 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | LN | LN | Score | P-Value |
|--|--|---|---|---|--|--|
| A17503001001_899177_c3_1335 | 3690 | 7462 | 930 | 309 | 596 | 5.2e-58 |
| Description | | | | | ., | |
| pir:[LN:G70046] [AC:G70046] [1] [OR:Bacillus subtilis] [DB:p: [AC:AJ223978] [PN:putative metal [OR:Bacillus subtilis] [DB:geng fragment from yvsA to yvqA.] [1] >gp:[GI:e1184397:g2635815] [LN [FN:unknown] [OR:Bacillus subticomplete genome (section 17 of iron-binding protein] [LE:20559] >gp:[GI:e1186006:g2635831] [LN [FN:unknown] [OR:Bacillus subticomplete genome (section 18 of iron-binding protein] [LE:3001] | ir2] >gp al bindi pept-bct LE:23792 :BSUB001 ilis] [D 21): fr 51] [RE: :BSUB001 ilis] [D 21): fr | :[GI:e1 ng prot 1] [DE:] [RE:2 7] [AC: B:genpe om 3197 206495] 8] [AC: B:genpe om 3399 | 249807: cein, Yv Bacillu 24736] 2299120: cpt-bctl 7001to 3 [DI:co 299121: cpt-bctl | g28328 yrC] [0 is subt [DI:din :AL0091] [DE: 3414420 pmpleme :AL0091] [DE: | B11] [LN SN:yvrC] cilis 42 cect] L26] [GN Bacillu D.] [NT: ent] L26] [GN | ::BS43KBDNA] ::7kB DNA ::yvrC] :s subtilis similar to ::yvrC] :s subtilis |
| ORF Name AI7503001001_900256_c1_1017 Description NO-HIT | NT ID | AA ID | NT LN 210 | AA LN 69 | <u>Score</u> | P-Value |
| ORF Name AI7503001001_959437_c1_965 Description NO-HIT | NT ID | <u>AA ID</u> 7464 | <u>NT</u> <u>LN</u> 159 | <u>AA</u> <u>LN</u> 52 | <u>Score</u> | P-Value |
| ORF Name AI7503001001_9642_f1_227 | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> 876 | <u>AA</u> <u>LN</u> 291 | <u>Score</u> | <u>P-Value</u> 2.4e-76 |
| Description pir:[LN:E70006] [AC:E70006] [I (undecapreno) yubB] [GN:yubB] protein bacA] [OR:Bacillus subt [LN:BSUB0016] [AC:Z99119:AL0091 | [CL:Esc cilis] [| herichi DB:pir2 | .a coli :] >gp:[| bacitr GI:e11 | acin re .85988:g | sistance 2635599] |

subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [NT:similar to bacitracin resistance

protein] [LE:195935] [RE:196765] [DI:complement]

| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|---|--|---|--------------------------------------|--|-----------------------------|
| A17503001001_969075_c1_1109 | 3694 | 7466 | 354 | 117 | 359 | 6.7e-33 |
| Description | · / L | -16 | · · · · · · · · · · · · · · · · · · · | l L | | |
| <pre>pir:[LN:B70021] [AC:B70021] [F [CL:hypothetical protein yjbD] >gp:[GI:e1184359:g2635777] [LN: [FN:unknown] [OR:Bacillus subticomplete genome (section 17 of arsenate reductase] [LE:168632]</pre> | [OR:Bac BSUB001 llis] [I 21): fi | cillus s 17] [AC: DB:genpe com 3197 | ubtili: Z99120 pt-bct: 001to : | s] [DB :AL009 1] [DE 341442 | :pir2] 126] [GN :Bacillu 0.] [NT: | :yusI] s subtilis |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_969555_f1_218 Description | 3695 | 7467 | 168 | 55 | J | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_969812_f1_63 | 3696 | 7468 | 123 | 40 | | • |
| Description NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503001001_970327_c2_1301 | 3697 | 7469 | 297 | 98 | 167 | 1.5e-12 |
| Description pir:[LN:F70020] [AC:F70020] [F [OR:Bacillus subtilis] [DB:pir2 [AC:Z99120:AL009126] [GN:yusE] [DB:genpept-bct1] [DE:Bacillus from 3197001to 3414420.] [NT:si [DI:complement] |] >gp: [FN:unk subtili | [GI:e118 known] [ks compl | 4355:g2 OR:Bac ete ger | 263577 illus nome (| 3] [LN:B subtilis section | SUB0017]] 17 of 21): |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503001001_973400_f3_690 | 3698 | 7470 | 180 | 59 | 43 | 0.013 |
| Description gp:[GI:e1363550:g4127809] [LN:E protein] [GN:orf6] [OR:Enteroco [DE:Enterococcus faecalis plasm [RE:4732] [DI:direct] | ccus fa | ecalis] | [DB:ge | enpept | -bct1] | |

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value | |
|---|---------------------|---------------|----------------|-----------------|----------|--------------|--|
| AI7503001001_978426_c1_1043 | 3699 | 7471 | 978 | 325 | 1338 | 1.2e-136 | |
| Description gp:[GI:e1330452:g3724155] [LN: | 52200535 | ים וארי | A.TOO53 | ים] וכב | V.membra | ene proteinl | |
| [GN:sstA] [FN:iron transport p: [DB:genpept-bct1] [DE:Staphylooperon.] [LE:273] [RE:1245] [DE: | rotein] coccus a | [OR:Staureus, | phyloc | occus a | aureus] | - | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
| A17503001001_9882950_c2_1204 Description | 3700 | 7472 | 1350 | 449 | 1090 | 2.3e-110 | |
| sp:[LN:YHDP_BACSU] [AC:O07585] [GN:YHDP] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 49.9 KD PROTEIN IN CITA-SSPB INTERGENIC REGION] [SP:O07585] [DB:swissprot] >pir:[LN:F69826] [AC:F69826] [PN:hemolysin homolog yhdP] [GN:yhdP] [CL:hypothetical protein HI0107] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182956:g2633290] [LN:BSUB0006] [AC:Z99109:AL009126] [GN:yhdP] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [NT:similar to hemolysin] [SP:007585] [LE:32043] [RE:33377] [DI:complement] >gp:[GI:e1191881:g2226211] [LN:BSY14082] [AC:Y14082] [PN:hypothetical protein] [GN:yhdP] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 72 to 75 degrees: spoVRto sspB.] [NT:Similarity to yhdT, this submission, and to] [SP:007585] [LE:18691] [RE:20025] [DI:complement] | | | | | | | |
| ORF Name | NT_ID | AA ID | NT LN | AA LN | Score | P-Value | |
| A17503001001_9884625_£2_464 | 3701 | 7473 | 1509 | 502 | 1226 | 9.0e-125 | |
| Description sp:[LN:YCLF_BACSU] [AC:P94408] [GN:YCLF] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 53.3 KD PROTEIN IN SFP-GERKA INTERGENIC REGION] [SP:P94408] [DB:swissprot] >pir:[LN:C69762] [AC:C69762] [PN:di-tripeptide ABC transporter (membrane pr) homolog yclF] [GN:yclF] [CL:peptide transporter protein] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182334:g2632668] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:yclF] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:similar to di-tripeptide ABC transporter | | | | | | | |

(membrane) [SP:P94408] [LE:13065] [RE:14543] [DI:complement] >gp:[GI:d1009635:g1805438] [LN:D50453] [AC:D50453] [PN:homologue of Di-tripeptide transporter Dtp of L.] [GN:yclF] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168 trpC2) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for 25-36 degree region containing theamyE-srfA region, complete cds.] [LE:95468] [RE:96946] [DI:complement]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
|--|---|---|--------------------------------------|--|--|---|--|
| A17503001001_989010_c1_1128 | 3702 | 7474 | 408 | 135 | 371 | 3.6e-34 | |
| Description | | | | | | | |
| pir:[LN:G70024] [AC:G70024] [IGN:yutM] [CL:conserved hypother [DB:pir2] >gp:[GI:e1184295:g263 [GN:yutM] [FN:unknown] [OR:Bacsubtilis complete genome (section in the section of the sectio | netical 35713] [illus su ion 17 o | protein LN:BSUB btilis] f 21): | HI0376 0017] [DB:ge from 31 | 5] [OR: [AC:Z99 enpept- 197001t | Bacillu 9120:ALC bct1] co 34144 | us subtilis] 009126] [DE:Bacillus | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
| AI7503012391_11291_c2_183 | 3703 | 7475 | 153 | 50 |] | | |
| Description | | | | | | | |
| NO-HIT | | | | | | | |
| ORF Name AI7503012391_1369012_c2_207 Description NO-HIT | <u>NT ID</u> | <u>AA ID</u> 7476 | NT LN | AA LN 41 | Score | P-Value | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | |
| AI7503012391_13790943_c1_161 | 3705 | 7477 | 1644 | 547 | 2797 | 3.0e-291 | |
| Description pir:[LN:C56976] [AC:C56976] [PN:transfer complex protein TrsK] [GN:trsK] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310618] [LN:STATRSC] [AC:L11998] [GN:trsK] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (individual_isolate pG01) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus conjugative transfer gene complex (trs).] [NT:putative] [LE:10475] [RE:12115] [DI:direct] >gp:[GI:g3676445] [LN:AF051917] [AC:AF051917:L19570] [PN:putative membrane protein TraK] [GN:traK] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [NT:similar to transfer-associated proteins of] [LE:34078] [RE:35718] [DI:direct] | | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
|---|----------------------------------|----------------------------------|----------------------------|----------------------------|-----------------------------|----------------|--|--|
| A17503012391_14507827_c3_220 | 3706 | 7478 | 168 | 55 | 273 | 8.7e-24 | | |
| Description | | | | - | | | | |
| pir:[LN:F56976] [AC:F56976] [[OR:Staphylococcus aureus] [DB [AC:AF051917:L19570] [PN:unkno [DB:genpept-bct2] [DE:Staphylo sequence.] [NT:Orf55; possibly [RE:37346] [DI:direct] | :pir2] : wn] [OR: coccus & | >gp:[GI: :Staphyl aureus p | g36764 ococcu lasmid | 48] [Li s aure pSK41 | N:AF0519 us] , comple | ete | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503012391_14844187_c2_195 | 3707 | 7479 | 3081 | 1026 | 5411 | 0.0 | | |
| Description | | | | | | | | |
| sp:[LN:SYIP_STAAU] [AC:P41368] [GN:MUPR] [OR:STAPHYLOCOCCUS AUREUS] [EC:6.1.1.5] [DE:(ISOLEUCINETRNA LIGASE) (ILERS) (MUPIROCIN RESISTANCE PROTEIN)] [SP:P41368] [DB:swissprot] >gp:[GI:g581558] [LN:SADNAMUPR] [AC:X75439] [PN:isoleucyl tRNA synthetase] [GN:ileS] [OR:Staphylococcus aureus] [DB:genpept-bctl] [DE:S.aureus plasmid encoded DNA, mup R gene.] [SP:P41368] [LE:477] [RE:3551] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> | | |
| AI7503012391_15631901_c3_234 | 3708 | 7480 | 645 | 214 | 943 | 8.8e-95 | | |
| Description | | | | | | | | |
| <pre>gp:[GI:g3676416] [LN:AF051917] [AC:AF051917:L19570] [PN:putative resolvase Res] [GN:res] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:2779] [RE:3336] [DI:direct]</pre> | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503012391_156910_f2_62 | 3709 | 7481 | 483 | 160 | 729 | 4.2e-72 | | |
| Description | | | | · - | | | | |
| gp:[GI:g3676415] [LN:AF051917] [AC:AF051917:L19570] [PN:unknown] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [NT:Orf149] [LE:2068] [RE:2484] [DI:complement] | | | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|---|----------------------|-------------------|----------|----------|----------|----------------|
| AI7503012391_19625062_c3_225 | 3710 | 7482 | 165 | 54 | 7 | |
| Description | | | | | J | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503012391_19665885_c1_175 | 3711 | 7483 | 1728 | 575 | 2999 | 0.0 |
| Description | | | | | | |
| <pre>gp:[GI:g3676419] [LN:AF051917] [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequence pRS01 LtrC] [LE:6350] [RE:8074]</pre> | :genpept ce.] [NT | -bct2] :Orf575 | [DE:Sta | aphylo | coccus a | ureus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_19744010_c3_232 | 3712 | 7484 | 288 | 95 | 466 | 3.1e-44 |
| <pre>gp:[GI:g3676414] [LN:AF051917] [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequence [DI:direct]</pre> | :genpept | -bct2] | [DE:Sta | phylo | coccus a | ureus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_2115625_f2_61 | 3713 | 7485 | 195 | 64 |] | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_2150037_c1_180 | 3714 | 7486 | 336 | 111 | 464 | 5.0e-44 |
| Description | | | | | | |
| <pre>gp:[GI:g3676421] [LN:AF051917] [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequence</pre> | genpept | -bct2] | [DE:Sta | phyloc | occus a | |

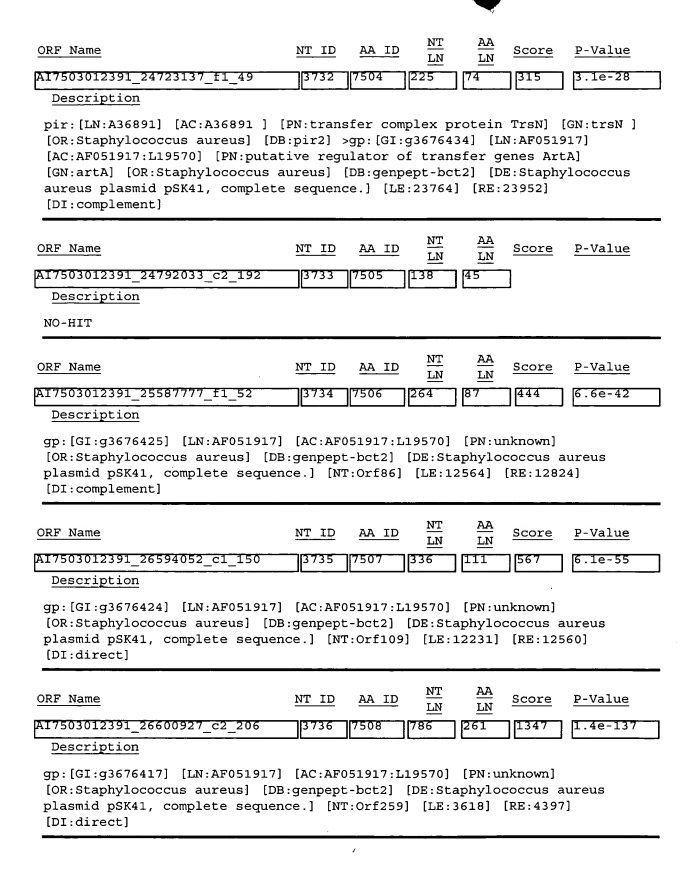
| ORF Name | NT ID | AA ID | $\frac{\text{NT}}{\text{LN}}$ | AA LN | Score | P-Value |
|---|--|--|--|-------------------------------------|--|----------------------|
| AI7503012391_21754035_f3_110 | 3715 | 7487 | 141 | 46 | ך | |
| Description | <u> </u> | JL | | | _ | |
| NO-HIT | | - - | _ | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_23468753_c3_236 | 3716 | 7488 | 1647 | 548 | 2838 | 1.4e-295 |
| Description | | | | | | |
| <pre>gp:[GI:g3676418] [LN:AF051917] [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequence [DI:direct]</pre> | :genpept | -bct2] | [DE:Sta | aphylo | coccus a | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_23594712_c2_211 | 3717 | 7489 | 309 | 102 | 392 | 2.1e-36 |
| Description | | | | | | |
| <pre>gp:[GI:g3676422] [LN:AF051917] protein] [OR:Staphylococcus au aureus plasmid pSK41, complete [DI:direct]</pre> | reus] [I | B:genpe | pt-bct2 | 2] [DE | :Staphyl | lococcus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_23603382_c2_196 | 3718 | 7490 | 603 | 200 | 688 | 9.2e-68 |
| Description sp:[LN:YIL2_STAAU] [AC:P41370] PROTEIN IN ILES 3'REGION (ORF (Orgin: [LN:S40262] [AC:S40262]] aureus] [DB:pir2] >gp:[GI:g4382] [OR:Staphylococcus aureus] [DB:DNA, mup R gene.] [NT:ORF C] [S | C) (FRAG [PN:hypc 228] [LN :genpept | MENT)] thetica :SADNAM -bct1] | [SP:P41 l prote UPR] [A [DE:S.a | l370] ein C] AC:X75 aureus | [DB:swis [OR:Sta 439] plasmid | ssprot] uphylococcus |

| ORF Name | NT ID | AA ID | $\frac{	ext{NT}}{	ext{LN}}$ | AA LN | Score | P-Value |
|---|--|--|---|---|---|--|
| AI7503012391_23651712_c2_191 | 3719 | 7491 | 480 | 159 | 779 | 2.1e-77 |
| Description pir: [LN:B56976] [AC:B56976] [[OR:Staphylococcus aureus] [DB [AC:L11998] [GN:trsJ] [OR:Stap (individual_isolate pG01) DNA] conjugative transfer gene comp [RE:10478] [DI:direct] >gp:[GI [PN:putative membrane protein [DB:genpept-bct2] [DE:Staphylo sequence.] [LE:33617] [RE:3408] | s:pir2] > hylococo [DB:gen blex (trs :g367644 TraJ] [G | egp:[GI: cus aure pept-bc despendent despend | tein cog31061'us] [SI t1] [DI :putat: AF0519: | omplex 7] [LN R:Stap E:Stap ive] [17] [A | :STATRS(hylococo hylococo LE:10014 C:AF0519 ococcus | C] cus aureus cus aureus 4] 917:L19570] aureus] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_23727302_c3_215 | 3720 | 7492 | 1296 | 431 | 2175 | 2.5e-225 |
| Description pir:[LN:G36891] [AC:G36891] [[OR:Staphylococcus aureus] [DB [AC:L11998] [GN:trsF] [OR:Stap (individual_isolate pG01) DNA] conjugative transfer gene comp [DI:direct] >gp:[GI:g3676440] [PN:putative membrane protein [DB:genpept-bct2] [DE:Staphylosequence.] [LE:28614] [RE:2989 | hylocococococococococococococococococococ | gp:[GI: us aure pept-bc].] [NT [1917] [N:traF] | g310613 us] [SI t1] [DI :putat: AC:AF05 [OR:St | 3] [LN R:Stap E:Stap ive] [51917: taphyle | :STATRSC hylococc hylococc LE:5011] L19570] ococcus | cus aureus cus aureus [[RE:6291] aureus] |
| ORF Name AI7503012391_23959802_c2_201 Description | NT ID | <u>AA ID</u> 7493 | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> 77 | Score | <u>P-Value</u> |
| NO-HIT | | | | | | |
| ORF Name AI7503012391_23959802_f1_16 | NT ID | <u>AA ID</u> | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> | Score | <u>P-Value</u> |
| Description | | | | | _ | |
| NO-HIT | | | | | | |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|------------------------------|-------|-------|----------|-----------------|-------|----------------|
| A17503012391_23959802_f1_32 | 3723 | 7495 | 135 | 44 |] | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | <u>P-Value</u> |
| AI7503012391_23959802_f1_51 | 3724 | 7496 | 123 | 40 |] | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_23959802_f2_70 | 3725 | 7497 | 174 | 57 |] | |
| Description | | | | | | |
| NO-HIT | | | ··· | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_23959802_f3_119 | 3726 | 7498 | 123 | 40 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_24001537_c2_187 | 3727 | 7499 | 750 | 249 | 1184 | 2.5e-120 |
| Description | | | | | | |

pir:[LN:E36891] [AC:E36891] [PN:transfer complex protein TrsD] [GN:trsD] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310611] [LN:STATRSC] [AC:L11998] [GN:trsD] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (individual_isolate pG01) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus conjugative transfer gene complex (trs).] [NT:putative] [LE:2283] [RE:2966] [DI:direct] >gp:[GI:g3676438] [LN:AF051917] [AC:AF051917:L19570] [PN:TraD] [GN:traD] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:25886] [RE:26569] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|---|--|---|---|---|---|
| AI7503012391_24298387_c1_171 | 3728 | 7500 | 147 | 48 | 68 | 0.045 |
| Description | | | | | | |
| sp:[LN:PSBH_CHLVU] [AC:P56323] [DE:PHOTOSYSTEM II 10 KD PHOSP | | | CHLOREI P56323] | | LGARIS] swisspro | ot] |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503012391_24393803_f1_28 | 3729 | 7501 | 171 | 56 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_24415937_c2_189 | 3730 | 7502 | 489 | 162 | 819 | 1.2e-81 |
| Description | | | | | | |
| pir:[LN:I36891] [AC:I36891] [I [OR:Staphylococcus aureus] [DB: [AC:L11998] [GN:trsH] [OR:Staph (individual_isolate pG01) DNA] conjugative transfer gene compl [DI:direct] >gp:[GI:g3676442] [PN:lipoprotein TraH] [GN:traH] [DE:Staphylococcus aureus plasm peptide recognized as a pheromo | pir2] > nylococo [DB:gen Lex (trs [LN:AF05] [OR:St nid pSK4 | gp:[GI:grays aurevalue aur | 3310615 us] [SF c1] [DE cputati AC:AF05 occus a lete se | E:Stap E:Stap E:Stap ve] [E1917: ureus equence | :STATRSC hylococc hylococc LE:7395] L19570]] [DB:ge e.] [NT: | eus aureus eus aureus [RE:7880] enpept-bct2] signal |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_24664812_c1_155 | 3731 | 7503 | 423 | 140 | 671 | 5.8e-66 |
| Description | | | | | | |
| pir:[LN:D36891] [AC:D36891] [EV:D36891] [E | pir2] > nylococc [DB:gen ex (trs [LN:AF05 TraC] [G | gp:[GI:gus aureupept-bct].] [NT:1917] [FN:traC]ureus pl | 3310610 is] [SR :1] [DE :putati AC:AF05 | [LN :Stapl :Stapl ve] [1 [1917:1] | :STATRSC hylococc hylococc LE:1889] L19570] ococcus | us aureus us aureus [RE:2296] |



| ORF Name AI7503012391_29558262_c2_205 Description | NT ID | <u>AA ID</u> 7509 | <u>NT</u> <u>LN</u> 129 | AA LN 42 | Score | P-Value | | |
|--|----------|----------------------|-------------------------------|----------------|----------|----------|--|--|
| NO-HIT | <u>.</u> | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503012391_30273550_c1_162 Description | 3738 | 7510 | 432 | 143 | 671 | 5.8e-66 | | |
| pir:[LN:E56976] [AC:E56976] [PN:transfer complex protein TrsM] [GN:trsM] [CL:single-stranded DNA-binding protein homology] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310620] [LN:STATRSC] [AC:L11998] [GN:trsM] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (individual_isolate pG01) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus conjugative transfer gene complex (trs).] [NT:putative] [LE:13127] [RE:13519] [DI:direct] >gp:[GI:g3676447] [LN:AF051917] [AC:AF051917:L19570] [PN:putative single-stranded DNA binding protein] [GN:traM] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:36730] [RE:37122] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503012391_30503392_c2_188 | 3739 | 7511 | 1083 | 360 | 1889 | 5.0e-195 | | |
| Description gp:[GI:g3676441] [LN:AF051917] protein TraG] [GN:traG] [OR:Sta [DE:Staphylococcus aureus plasm [RE:30988] [DI:direct] | phyloco | ccus au | reus] | [DB:gei | npept-bo | t2] | | |

NT AΑ NT ID AA ID Score P-Value ORF Name LNLN AI7503012391 33786251 c3 230 3740 7512 1113 370 1926 6.0e-199

Description

sp:[LN:AACA STAAU] [AC:P14507] [GN:AACA-APHD] [OR:STAPHYLOCOCCUS AUREUS: ENTEROCOCCUS FAECALIS] [SR:, STREPTOCOCCUS FAECALIS] [EC:2.3.1.-:2.7.1.-] [DE:AMINOGLYCOSIDE PHOSPHOTRANSFERASE, (APH(2''))] [SP:P14507] [DB:swissprot] >pir:[LN:S26353] [AC:S26353] [PN:aminoglycoside resistance protein aacA-aphD] [OR:Staphylococcus aureus] [DB:pir2] >pir:[LN:A26048] [AC:A26048] [PN:aminoglycoside acetyltransferase] [OR:Enterococcus faecalis] [DB:pir2] >gp:[GI:g152948] [LN:STAAGLSRA] [AC:M18086:M29261] [PN:aminoglycoside resistance protein] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (strain SK982) DNA] [DB:genpept-bct1] [DE:S.aureus transposon 4001 aacA-aphD aminoglycoside resistance gene, complete cds, and right and left IS256 transposase genes.] [LE:1725] [RE:3164] [DI:direct] >gp:[GI:g153586] [LN:STRBRP] [AC:M13771] [OR:Enterococcus faecalis] [SR:S.faecalis DNA, clone pSF815A] [DB:genpept-bct1] [DE:Streptococcus faecalis 6'-aminoglycoside acetyltransferasephosphotransferase (AAC(6')-APH(2')) bifunctional resistanceprotein, complete cds.] [NT:AAC(6')-APH(2') bifunctional resistance protein] [LE:304] [RE:1743] [DI:direct] >gp:[GI:g3676454] [LN:AF051917] [AC:AF051917:L19570] [PN:bifunctional aminoglycoside modifying enzyme] [GN:aacA-aphD] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:42505] [RE:43944] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|-----------------------------|-------|-------|----------|----------|-------|---------|
| AI7503012391_33786251_f2_66 | 3741 | 7513 | 390 | 129 | 644 | 4.2e-63 |

Description

sp:[LN:TRA6_STAAU] [AC:P19775] [GN:TNP] [OR:STAPHYLOCOCCUS AUREUS]
[DE:TRANSPOSASE FOR INSERTION SEQUENCE ELEMENT IS256 IN TRANSPOSON TN4001]
[SP:P19775] [DB:swissprot] >pir:[LN:JS0296] [AC:JS0296] [PN:transposase]
[GN:tnp] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g152947]
[LN:STAAGLSRA] [AC:M18086:M29261] [PN:transposase] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (strain SK982) DNA] [DB:genpept-bct1]
[DE:S.aureus transposon 4001 aacA-aphD aminoglycoside resistance
gene,complete cds, and right and left IS256 transposase genes.] [LE:102]
[RE:1274] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
|--|---|---|--|--|--|-------------------------------------|--|--|
| AI7503012391_34242300_c1_158 | 3742 | 7514 | 2037 | 678 | 3430 | 0.0 | | |
| Description | | J | | | | | | |
| pir:[LN:F36891] [AC:F36891] [B [OR:Staphylococcus aureus] [DB [AC:L11998] [GN:trsE] [OR:Staph (individual_isolate pG01) DNA] conjugative transfer gene comp [DI:direct] >gp:[GI:g3676439] [PN:putative ATPase TraE] [GN:60] [DB:genpept-bct2] [DE:Staphylocosequence.] [LE:26584] [RE:2860] | :pir2] > hylococc [DB:gen lex (trs [LN:AF05 traE] [C coccus a | gp:[GI: us aure pept-bc).] [NT [1917] [OR:Staph ureus p | g310612 us] [SI t1] [DI :putat: AC:AF0! ylococo | 2] [LN R:Stap E:Stap ive] [51917: cus au | :STATRSC hylococc hylococc LE:2981] L19570] reus] | us aureus us aureus [RE:4999] | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| A17503012391_34642562_c1_174 | 3743 | 7515 | 207 | 68 | | | | |
| Description | | | | | | | | |
| NO-HIT | | | | | | | | |
| | - | | | - | | | | |
| ORF Name | NT ID | AA ID | $\frac{\mathtt{NT}}{\mathtt{LN}}$ | <u>AA</u> LN | Score | <u>P-Value</u> | | |
| AI7503012391_35397177_c2_182 | 3744 | 7516 | 1062 | 353 | 1760 | 2.3e-181 | | |
| Description | | | | | | | | |
| gp:[GI:g3676423] [LN:AF051917] [AC:AF051917:L19570] [PN:unknown] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [NT:Orf346] [LE:11188] [RE:12228] [DI:direct] | | | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value | | |
| AI7503012391_35397177_c3_241 | 3745 | 7517 | 132 | 44 | 142 | 3.1e-09 | | |
| Description | | | | | | | | |
| gp:[GI:g3676423] [LN:AF051917] [AC:AF051917:L19570] [PN:unknown] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [NT:Orf346] [LE:11188] [RE:12228] [DI:direct] | | | | | | | | |

| ORF Name | NT ID | AA ID | <u>NT</u> LN | <u>AA</u> LN | Score | <u>P-Value</u> |
|--|--|--|--|-------------------------------------|--|--------------------------------|
| A17503012391_36048212_c3_221 | 3746 | 7518 | 774 | 257 | 783 | 7.9e-78 |
| Description | | | | | | |
| sp:[LN:YIL1_STAAU] [AC:P41369] PROTEIN IN ILES 5'REGION (ORF >pir:[LN:S40261] [AC:S40261] aureus] [DB:pir2] >gp:[GI:g438 [OR:Staphylococcus aureus] [DB DNA, mup R gene.] [NT:ORF B] [| B) (FRAC [PN:hypo 227] [LN :genpept | SMENT)] othetica J:SADNAM :-bct1] | [SP:P4] l prote [UPR] [[DE:S.a | 1369] ein B] AC:X75 aureus | [DB:swis [OR:Sta 439] plasmid | ssprot] aphylococcus d encoded |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503012391_36605260_c1_169 | 3747 | 7519 | 630 | 209 | 1077 | 5.5e-109 |
| Description | | | | | | |
| <pre>gp:[GI:g3676413] [LN:AF051917] [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequen</pre> | :genpept | -bct2] | [DE:Sta | aphylo | coccus a | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_3907943_f3_103 | 3748 | 7520 | 2010 | 669 | 3498 | 0.0 |
| Description | | | | | | |
| <pre>gp:[GI:g3676420] [LN:AF051917] [GN:nes] [OR:Staphylococcus au aureus plasmid pSK41, complete [DI:complement]</pre> | reus] [I | B:genpe | pt-bct2 | 2] [DE | :Staphyl | _ |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_3908407_c3_214 | 3749 | 7521 | 123 | 40 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name AI7503012391_4300002_c1_153 Description | NT ID | <u>AA ID</u> 7522 | <u>NT</u> <u>LN</u> | <u>AA</u> <u>LN</u> 41 | Score | P-Value |
| NO-HIT | | | | | | |
| | | | | | | • |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|--|--|---|---|--|---|
| AI7503012391_4428802_f2_57 | 3751 | 7523 | 168 | 55 | 7 | |
| Description | 1L | JL1 | | L | J | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503012391_4486075_c1_154 Description | 3752 | 7524 | 975 | 324 | 1640 | 1.2e-168 |
| pir:[LN:B36891] [AC:B36891] [OR:Staphylococcus aureus] [D [AC:L11998] [GN:trsA] [OR:Sta (individual_isolate pG01) DNA conjugative transfer gene com [DI:direct] >gp:[GI:g3676435] [GN:traA] [OR:Staphylococcus aureus plasmid pSK41, complet | B:pir2] > phylococo] [DB:gen plex (trs [LN:AF05 aureus] [| gp:[GI:g us aureu pept-bct).] [NT: 1917] [A DB:genpe | 310608 [SR] [SR 1] [DE putati .C:AF05 | [LN:Stap ::Stap ve] [1917: 2] [D | :STATRSC hylococo hylococo LE:521] L19570] E:Staphy | cus aureus cus aureus [RE:1495] [PN:TraA] vlococcus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_4586062_c1_176 | 3753 | 7525 | 141 | 46 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| A17503012391_4724035_c1_170 | 3754 | 7526 | 1134 | 377 | 1728 | 5.7e-178 |
| Description | | | | • | | |
| <pre>gp:[GI:g3676414] [LN:AF051917 [OR:Staphylococcus aureus] [DI plasmid pSK41, complete seque: [DI:direct]</pre> | B:genpept | -bct2] [| DE:Sta | phylo | coccus a | |

NT AA ORF Name NT ID AA ID Score P-Value LNLN 705 AI7503012391 4745437 c2 190 3755 7527 2118 3603 0.0 Description

pir:[LN:A56976] [AC:A56976] [PN:transfer complex protein TrsI] [GN:trsI] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310616] [LN:STATRSC] [AC:L11998] [GN:trsI] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (individual_isolate pG01) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus conjugative transfer gene complex (trs).] [NT:putative] [LE:7896] [RE:9998] [DI:direct] >gp:[GI:g3676443] [LN:AF051917] [AC:AF051917:L19570] [PN:putative topoisomerase TraI] [GN:traI] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:31499] [RE:33601] [DI:direct]

NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503012391_5109675_c2_186 107 3756 7528 324 535 1.5e-51

Description

pir:[LN:C36891] [AC:C36891] [PN:transfer complex protein TrsB] [GN:trsB] [OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310609] [LN:STATRSC] [AC:L11998] [GN:trsB] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (individual_isolate pG01) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus conjugative transfer gene complex (trs).] [NT:putative] [LE:1512] [RE:1829] [DI:direct] >gp:[GI:g3676436] [LN:AF051917] [AC:AF051917:L19570] [PN:putative membrane protein TraB] [GN:traB] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus plasmid pSK41, complete sequence.] [LE:25115] [RE:25432] [DI:direct]

NTAΑ ORF Name NT ID AA ID Score P-Value LN LN AI7503012391_5314202_c3_219 3757 7529 918 305 1524 2.4e-156

Description

pir:[LN:D56976] [AC:D56976] [PN:transfer complex protein TrsL] [GN:trsL]
[OR:Staphylococcus aureus] [DB:pir2] >gp:[GI:g310619] [LN:STATRSC]
[AC:L11998] [GN:trsL] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus
(individual_isolate pG01) DNA] [DB:genpept-bctl] [DE:Staphylococcus aureus
conjugative transfer gene complex (trs).] [NT:putative] [LE:12193]
[RE:13110] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|---|---|--|---|--|---|
| A17503012391_6929686_c1_152 | 3758 | 7530 | 693 | 230 | 1199 | 6.5e-122 |
| Description | <u> </u> | | | | | |
| gp:[GI:g1762100] [LN:SEU40385] [OR:Staphylococcus epidermidis epidermidis plasmid pSK818 insetransposase gene, complete cds [RE:731] [DI:direct] >gp:[GI:g:[PN:transposase] [GN:tnp] [OR:EDE:Staphyloccous epidermidis putative transposase gene, complete:57] [RE:731] [DI:direct] |] [DB:go ertion : .] [NT:' 1762102] Staphylo plasmid | enpept-bosequencel Inp; puta [LN:SEU pcoccus e pSK818 i | ct1] [I [S257(8 ative t J40386] epidern | DE:Sta 818B) transp [AC: midis] ion se | phylocco putative osase] [U40386] [DB:gen quenceIS | pus [LE:57] [pept-bct1] [257(818C) |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012391_6929686_c1_163 | 3759 | 7531 | 693 | 230 | 1201 | 4.0e-122 |
| Description | · | | | | | |
| pir: [LN:A60634] [AC:A60634:C306] [OR:Staphylococcus aureus] [DB [AC:X53952] [PN:transposase] [OE:S.aureus plasmid pSH6 DNA [LE:188] [RE:862] [DI:direct] [AC:AF051917:L19570] [PN:putate [OR:Staphylococcus aureus] [DB plasmid pSK41, complete sequence | :pir2] : OR:Stapl for inse >gp:[GI ive tran :genpept | >gp:[GI:g nylococcu ertion se :g3676452 nsposase c-bct2] [| y46597] us aure equence e] [LN: TnpE] [DE:Sta | [LN: eus] [es IS2 :AF051 [GN:t aphylo | SAIS2571 DB:genpe 57-1 and 917] npE] coccus a | ept-bct1] [IS256.] [ureus |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503012391_6929686_c1_168 | 3760 | 7532 | 693 | 230 | 1205 | 1.5e-122 |
| <u>Description</u> | | | | | | |
| pir:[LN:C60634] [AC:C60634:S26: [OR:Staphylococcus aureus] [DB [AC:X53951] [PN:putative transpose; [DB:genpept-bct1] [DE:S.aureus] IS257-2, IS257-3and IS256.] [LN:Spp:[GI:g3676456] [LN:AF051917] transposase TnpG] [GN:tnpG] [OI:DE:Staphylococcus aureus plasmannes] [RE:46387] [DI:direct] | :pir2] : posase] plasmic E:1752] [AC:AI R:Staphy | >gp:[GI:g [OR:Stap d pSH6 DN [RE:2426 F051917:L /lococcus | y46600] hyloco JA for [] [DI: J19570] s aureu | [LN: ccus inser direc [PN:] | SAIS2572 aureus] tion seq t] putative B:genpep | uences t-bct2] |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
|--|---|---|--|---|--|---------------------------------------|
| A17503012391_6929686_c2_197 | 3761 | 7533 | 560 | 219 | 1116 | 4.1e-113 |
| Description | _ | | | | | |
| pir:[LN:A60634] [AC:A60634:C3 [OR:Staphylococcus aureus] [D [AC:X53952] [PN:transposase] [DE:S.aureus plasmid pSH6 DNA [LE:188] [RE:862] [DI:direct] [AC:AF051917:L19570] [PN:puta [OR:Staphylococcus aureus] [D plasmid pSK41, complete seque | B:pir2] > [OR:Staph for inse >gp:[GI: tive tran B:genpept | gp:[GI:g ylococcu rtion se g3676452 sposase -bct2] [| 46597] s aure quence] [LN: TnpE] DE:Sta | [LN: eus] [es IS2 :AF051 [GN:taphylo | SAIS2571 DB:genpe 57-1 and 917] npE] coccus a | .] ept-bct1] lIS256.] uureus |
| ORF_Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012391_6929686_c3_229 | 3762 | 7534 | 93 | 230 | 1201 | 4.0e-122 |
| Description | | | | | | |
| [OR:Staphylococcus aureus] [DE:X53952] [PN:transposase] [DE:S.aureus plasmid pSH6 DNA [LE:188] [RE:862] [DI:direct] [AC:AF051917:L19570] [PN:puta [OR:Staphylococcus aureus] [DE:Dasmid pSK41, complete seques | [OR:Staph for inse >gp:[GI: tive tran B:genpept | ylococcu rtion se g3676452 sposase -bct2] [| s aure quence] [LN: TnpE] DE:Sta | eus] [es IS2 :AF051 [GN:t aphylo | DB:genpe 57-1 and 917] npE] coccus a | ept-bct1] NS256.] Tureus |
| ORF Name | NT ID | <u>AA ID</u> | NT LN | AA LN | Score | P-Value |
| AI7503012391_6929686_f3_115 Description | 3 / 6 3 | 7535 | | 227 | 1136 | 1.46-121 |
| pir: [LN:B60634] [AC:B60634:S26] [OR:Staphylococcus aureus] [DR:Staphylococcus aureus] [DR:Gac:X53951] [PN:putative trans [DB:genpept-bct1] [DE:S.aureus [S257-2, IS257-3and IS256.] [Sygp:[GI:g3676455] [LN:AF051915] [GN:tnpF] [GN:tnpF] [GE:Staphylococcus aureus plas [RE:45182] [DI:complement] | B:pir2] > sposase] s plasmid LE:556] [7] [AC:AF OR:Staphy | gp:[GI:g [OR:Stap pSH6 DN RE:1221] 051917:L lococcus | 46599] hyloco A for [DI:co 19570] aureu | [LN: occus inser comple [PN: | SAIS2572 aureus] tion seq ment] putative B:genpep | quences st-bct2] |

| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
|--|-----------|-----------------------|----------|----------|-------|----------------|
| A17503012391_822802_c3_213 | 3764 | 7536 | 011 | 336 | 1647 | 2.2e-169 |
| Description | | | | | | |
| gp:[GI:g3676426] [LN:AF051917] initiation protein Rep] [GN:re [DB:genpept-bct2] [DE:Staphylosequence.] [LE:13205] [RE:1416 | ep] [OR:S | Staphyloc ureus pl | occus | aureu | s] | _ |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| A17503012392_1442583_c1_19 | 3765 | 7537 | .44 | 47 | | |
| Description | | | | | _ | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012392_1442583_c3_25 | 3766 | 7538 | 44 | 47 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012392_23671885_c1_18 | 3767 | 7539 | .59 | 52 | | |
| Description | | | | | | |
| NO-HIT | | | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | P-Value |
| AI7503012392_2776391_f3_12 | 3768 | 7540 | 207 | 68 |] | |
| Description | | | | | | |
| NO-HIT | | _ | | | | |
| ORF Name | NT ID | AA ID | NT LN | AA LN | Score | <u>P-Value</u> |
| AI7503012392_29323431_f3_15 Description | 3769 | 7541 | .23 | 40 | _ | |
| NO-HIT | | | | | | |





NT ORF Name NT ID AA ID Score P-Value LN LN 7542 AI7503012392 30495306 f2 6 123 40 3770 Description NO-HIT NT AAORF Name NT ID AA ID P-Value Score LN LN AI7503012392 34023375 c2 23 7543 166 790 1.4e-78 501

Description

sp:[LN:REMA STAAU] [AC:P13969] [GN:REPL:REP] [OR:STAPHYLOCOCCUS AUREUS: STAPHYLOCOCCUS SIMULANS] [DE: REPLICATION AND MAINTENANCE PROTEIN (PLASMID REPLICATION PROTEIN)] [SP:P13969] [DB:swissprot] >pir:[LN:A29827] [AC:A29827:C46568] [PN:replication protein REP] [OR:Staphylococcus aureus] [DB:pir2] >qp:[GI:q153064] [LN:PE5PE5A] [AC:M17990] [GN:repL] [OR:Plasmid pE5] [SR:Plasmid pE5 DNA] [DB:genpept-bct1] [DE:Plasmid pE5 (from Staphylococcus aureus, strain RN451) repL proteinand ermC protein, complete cds.] [LE:389] [RE:865] [DI:direct] >gp:[GI:g506624] [LN:STAPT48CG] [AC:M19652] [OR:Plasmid pT48] [SR:Plasmid pT48 (from S.aureus strain T48) DNA] [DB:qenpept-bct1] [DE:Plasmid pT48 (from S.aureus strain T48) complete genome.] [NT:putative. ORF B.] [LE:64] [RE:540] [DI:complement] >gp:[GI:g2407672] [LN:AF019140] [AC:AF019140] [PN:Rep] [GN:rep] [OR:Staphylococcus simulans] [DB:genpept-bct2] [DE:Staphylococcus simulans erythromycin resistance plasmid pPV142 rRNAN-6-methyltransferase (ermM) and replication protein (rep) genes, complete cds.] [NT:plasmid replication protein] [LE:1699] [RE:2175] [DI:direct] >qp:[GI:q1791222] [LN:SCU82607] [AC:U82607] [PN:plasmid replication protein] [OR:Staphylococcus chromogenes] [DB:genpept-bct2] [DE:Staphylococcus chromogenes plasmid pPV141 erythromycin resistanceplasmid, rRNA N-6-methyltransferase (ermM) and plasmid replicationprotein genes, complete cds.] [LE:1583] [RE:2059] [DI:direct]

| ORF Name | NT ID | AA ID | NT LN | <u>AA</u> LN | Score | P-Value |
|----------------------------|-------|-------|----------|-----------------|-------|----------|
| A17503012392_4304683_c2_21 | 3772 | 7544 | 771 | 256 | 1270 | 2.0e-129 |
| Doggwintion | | | | | | |

Description

sp:[LN:ERM4 STAAU] [AC:P13978] [GN:ERMC] [OR:STAPHYLOCOCCUS AUREUS] [EC:2.1.1.48] [DE:RESISTANCE PROTEIN)] [SP:P13978] [DB:swissprot]

>pir:[LN:B46568] [AC:B46568] [PN:ermC protein] [CL:rRNA

(adenine-N6-)-methyltransferase] [OR:Staphylococcus aureus] [DB:pir2]

>gp:[GI:g455358] [LN:STAPT48CG] [AC:M19652] [PN:23S RNA methylase] [GN:ermC]

[OR:Plasmid pT48] [SR:Plasmid pT48 (from S.aureus strain T48) DNA]

[DB:genpept-bct1] [DE:Plasmid pT48 (from S.aureus strain T48) complete

genome.] [LE:988] [RE:1722] [DI:complement]